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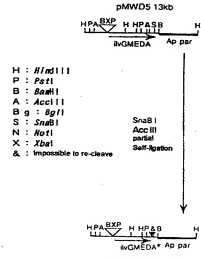
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# (54) PROCESS FOR PRODUCING L-VALINE AND L-LEUCINE

(57) L-valine is produced by culturing a microorganism beloging to the genus <u>Eschrichia</u> with the capability of producing L-valine or L-leucine wherein it requires lipoic acid for growth, a microorganism belonging to the genus <u>Eschrichia</u> with the capability of producing L-valine or L-leucine wherein it is deficient in H<sup>+</sup>-ATPase activity, a microorganism belonging to the genus <u>Eschrichia</u> with the capability of producing L-valine or L-leucine wherein it requires lipoic acid for growth and is deficient in H<sup>+</sup>-ATPase activity, in the liquid medium to allow the L-valine to be produced and accumulated in a culture medium, and collecting it.

FIG. 5



pMWdAR6 11.7kb

### Description

## Technical Field

This invention relates to a microorganism belonging to the genus <u>Escherichia</u> having the capability of producing L-valine or L-leucine and, more particularly, a microorganism whose capability of producing L-valine or L-leucine is enhanced.

#### Background Art

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In the past, L-valine and L-leucine have been produced by a method of fermentation primarily using a microorganism belonging to the genus <u>Brevibacterium</u>, <u>Corynebacterium</u> or <u>Serratia</u> or a mutant thereof which produces L-valine or L-leucine (Amino acid fermentation, JAPAN SCIENTIFIC SOCIETY'S PRESS, pp.397-422, 1986). Although the conventional methods have considerably enhanced the productivity of these amino acids, the development of a more efficient, cost-effective technique is required in order to meet increasing demand for L-valine and L-leucine in the future.

On the other hand, a microorganism belonging to the genus <u>Escherichia</u> is potentially utilized as a potent L-valine or L-leucine-producing microorganism due to its rapid growth rate, progress in genetic analysis and plentiful genetic materials. However, there are few reports documenting the production of these amino acids with from <u>Escherichia</u> microorganisms, and as for L-branched chain amino acids, only a few reports deal with the production of L-isoleucine (Japanese Patent Application Laid-Open No. 5-304969(1993) and Japanese Patent Application Laid-Open No. 5-130882(1993).

#### Disclosure of the Invention

The object of the present invention, in view of the aforementioned points, is to provide an efficient and cost-effective method for producing L-valine and L-leucine by enhancing the capability of producing L-valine or L-leucine of a microorganism belonging to the genus

### Escherichia.

As a result of a wholeheartedly conducted study of the production of L-valine and L-leucine by mutants of microorganisms belonging to the genus <u>Escherichia</u>, the present inventors have found that a mutation, whereby lipoic acid is required for growth and/or H<sup>+</sup>-ATPase is deficient, enhances the capability of producing L-valine or L-leucine of a L-valine or L-leucine-producing microorganism.

Thus, a first microorganism of the present invention is a microorganism belonging to the genus <u>Escherichia</u> and having the capability of producing L-valine or L-leucine, which requires lipoic acid for growth. A second microorganism of the present invention is a microorganism belonging to the genus <u>Escherichia</u> and having the capability of producing L-valine or L-leucine, which is deficient in H+-ATPase. Furthermore, a third microorganism of the present invention is a microorganism belonging to the genus <u>Escherichia</u> and having the capability of producing L-valine or L-leucine, which requires lipoic acid for growth and is deficient in H+-ATPase.

The present invention also provides a method for producing L-valine or L-leucine comprising culturing the aforementioned microorganism in a liquid medium to allow the L-valine or L-leucine to be produced and accumulated in the medium and collecting it.

In the specification, the phrase "H+-ATPase deficient" means that cells do not substantially express H+-ATPase activity, and includes both of that an H+-ATPase gene does not express due to entire or partial deletion of an atp operon coding for eight subunits of H+-ATPase or split of the atp operon and that the H+-ATPase gene has substitution, insertion or deletion of one or more nucleotides therein so that the H+-ATPase protein which is produced by expression of the gene does not have H+-ATPase activity. The <a href="ilvGMEDA">ilvGMEDA</a> operon means a operon including each of <a href="ilvG, ilvM">ilvG</a>, <a href="ilvGMEDA">ilvD</a> genes, and the operon may additionally include <a href="ilvA">ilvA</a> gene, which expresses inactivated threonine deaminase, or may not include <a href="ilvA">ilvA</a> gene substantially.

The invention will be described in detail as follows:

## (1) Microorganism of the Present Invention

A microorganism of the invention is one which belongs to the genus <u>Escherichia</u> and has the capability of producing L-valine or L-leucine and has any one of the following properties:

1. Lipoic acid is required for growth.

- 2. H+-ATPase is deficient.
- 3. Lipoic acid is required for growth and H+-ATPase activity is deficient.

In the present invention, the microorganism may possess any one of the aforementioned properties 1 to 3, and preferably possess property 3.

A microorganism having such properties can be obtained by giving the capability of producing L-valine or L-leucine to a microorganism belonging to the genus <u>Escherichia</u>, which is mutated so that it requires lipoic acid for growth and/or is deficient in H<sup>+</sup>-ATPase, or by enhancing the capability of producing L-valine or L-leucine in the aforementioned mutant. The microorganism of the present invention can be also obtained by inducing a mutation whereby lipoic acid is required for growth and/or a mutation whereby H<sup>+</sup>-ATPase is deficient in a microorganism belonging to the genus <u>Escherichia</u>.

The microorganism to be used in obtaining the aforementioned microorganisms can include a strain, which belongs to the genus <u>Escherichia</u> such as <u>Escherichia</u> coli (hereinafter, also referred to as <u>E. coli</u>) and exhibits no pathogenicity. For example, the following strains can be used.

Escherichia coli K-12 (ATCC10798) Escherichia coli W3110 (ATCC27325) Escherichia coli W1485 (ATCC12435)

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In order to introduce a mutation whereby lipoic acid is required for growth and/or a mutation whereby H<sup>+</sup>-ATPase is deficient into these microorganisms belonging to the genus <u>Escherichia</u>, the usual methods for introducing mutation, such as irradiation with X-ray or ultraviolet rays, or contact with mutagens including N-methyl-N'-nitro-N-nitrosoguanidine (hereinafter abbreviated as NG) and nitrous acid, can be applied. Additionally, the introduction of a mutation into a microorganism belonging to the genus <u>Escherichia</u> can be carried out by other genetic technique such as gene recombination, transduction, cell fusion and the like.

An example of the means for obtaining a mutant is as follows:

A mutant which requires lipoic acid for growth (hereinafter, referred to as a lipoic-acid-requiring strain) is obtained by culturing mutagenized bacterial cells on an agar plate, and by isolating colonies which exhibit lipoic acid requirement (A.A. Herbert and J.R. Guest: J. Gen. Microbiol., <u>53</u>, 363-381 (1968)). As a lipoic acid requiring strain, specifically, <u>E. coli</u> W1485lip2 (ATCC25645) can be used.

A mutant which is deficient in H\*-ATPase (hereinafter, referred to as a H\*-ATPase-deficient strain) is obtained by selecting mutants which cannot grow on an agar plate containing citric acid as the sole carbon source and can grow on an agar plate containing glucose as the sole carbon source from mutagenized bacterial cells, and by further selecting, from these mutants, strains which do not exhibit H\*-ATPase activity. As a H\*-ATPase-deficient strain, specifically, <u>E. coli</u> AN718 (E. coli Genetic Stock Center, Yale University, Department of Biology) can be used.

H<sup>+</sup>-ATPase is a membrane-binding enzyme with approximately 500,000 KD in molecular weight, in which 8 types of subunits complicatedly associate, and functions to pump H<sup>+</sup> outside of cytoplasm through changes in the free energy caused by ATP hydrolyzation and to synthesize ATP utilizing a H<sup>+</sup>-concentration gradient between the inside and outside of cytoplasmic membrane caused by intracellular respiration. Additionally, this enzyme is divided into an F0 fraction, which is localized on the inner membrane and exhibits H<sup>+</sup>-transport activity, and an F1 fraction, which is localized in the membrane surface and catalyzes the decomposition and synthesis of ATP, and the F0 is composed of 3 types of subunits a, b and c, while the F1 is composed of 5 types of subunits  $\alpha$ ,  $\beta$ ,  $\gamma$ ,  $\delta$ ,  $\epsilon$ . A strain which has a mutation in any of these subunits can be used as a H<sup>+</sup>-ATPase-deficient strain. The mutation of the H<sup>+</sup>-ATPase deficiency may include the expression of a mutant subunit, and the non-expression of subunits comprising H<sup>+</sup>-ATPase by the mutation at a promoter site.

Further, because oxidative phosphorylation is not carried out in a H<sup>+</sup>-ATPase-deficient strain and energy is obtained by substrate-level phosphorylation, it is expected that the addition of various agents including H<sup>+</sup>-ATPase inhibitors, TCA cycle inhibitors, respiratory chain inhibitors and uncoupling agents to the culture medium results in the same effect as H<sup>+</sup>-ATPase deficiency. Such H<sup>+</sup>-ATPase inhibitors include dicyclohexylcarbodiimide, tributyltin, and aurovertin, TCA cycle inhibitors include malonic acid, monoiodoacetic acid, methyl violet and 2,4-dinitrophenol, electron transport inhibitors include thenoyltrifluoroacetone, 2-n-nonyl-4-hydroxyquinoline-N-oxide and antimycin, and uncoupling agents include valinomycin, atebrin and 4,5,6,7-tetrafluoro-2-trifluoromethylbenzimidazol. These inhibitors may be used either alone or as a mixture of more than two types of inhibitors.

The lipoic-acid-requiring strain obtained as above is additionally mutagenized as a parent strain for selecting an H<sup>+</sup>-ATPase-deficient strain, or the H<sup>+</sup>-ATPase-deficient strain is additionally mutagenized as a parent strain for selecting a strain which comes to require lipoic acid, whereby a mutant which exhibits both lipoic acid requirement and H<sup>+</sup>-ATPase deficiency (hereinafter referred to as lipoic acid-requiring-H<sup>+</sup>-ATPase-deficient strain can be obtained. Further, the mutant which exhibits both lipoic acid requirement and H<sup>+</sup>-ATPase deficiency can be obtained by introducing one of

these mutations in a mutant which exhibits the other mutation by transduction, transformation, cell fusion and the like.

For example, lipoic acid-requiring-H\*-ATPase deficient strain can be obtained by transducing H\*-ATPase deficiency into a lipoic-acid-requiring strain as a parent strain. In this case, the aforementioned W1485lip2 strain can be used as a parent strain, and the aforementioned AN718 strain can be used as a donor strain. The lipoic acid-requiring-H\*-ATPase-deficient strain can be obtained by transducing a lipoic acid requirement into a H\*-ATPase-deficient strain as a parent strain.

The lipoic acid-requiring-H\*-ATPase-deficient strain can include <u>E. coli</u> AJ12631. The strain AJ12631 has been deposited on July 24, 1991 in National Institute of Bioscience and Human Technology of Agency of Industrial Science and Technology under an accession number of FERM P-12381, transferred to international deposition based on the Budapest Treaty on August 29, 1995, and deposited under an accession number of FERM BP-5209.

A microorganism of the present invention can be obtained by giving the capability of producing L-valine or L-leucine to a lipoic acid requiring mutant, a H+-ATPase-deficient mutant or a lipoic acid-requiring-H+-ATPase-deficient mutant belonging to the genus <u>Escherichia</u>, or by enhancing the capability of producing L-valine or L-leucine in the aforementioned mutants. Additionally, the microorganism of the present invention can be also obtained by introducing a lipoic acid requirement and/or H+-ATPase deficiency in a microorganism belonging to the genus <u>Escherichia</u> having the capability of producing L-valine or L-leucine. Further, even for a microorganism being low in the capability of producing L-valine or L-leucine, this capability can be enhanced by introducing a lipoic acid requirement and/or H+-ATPase deficiency.

#### (1) L-valine-producing microorganism

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An L-valine-producing microorganism can be obtained by giving the capability of producing L-valine to a lipoic-acid-requiring mutant, a H\*-ATPase-deficient mutant, or a lipoic acid-requiring-H\*-ATPase-deficient mutant of a microorganism belonging to the genus <u>Escherichia</u>, or by enhancing the capability of producing L-valine in the aforementioned mutant.

The giving or enhancing of the capability of producing L-valine is carried out, for example, by introducing genes for L-valine biosynthesis, whose regulatory mechanism is substantially released, into a microorganism belonging to the genus <u>Escherichia</u>. A mutation which leads to the suppression of the regulatory mechanism of the genes for the L-valine biosynthesis of the microorganism belonging to the genus <u>Escherichia</u>, may be introduced.

In a microorganism belonging to the genus <u>Escherichia</u>, the final step of L-valine biosynthesis is carried out by a group of <u>ilvGMEDA</u> operon-encoded enzymes. The <u>ilvGMEDA</u> operon includes each of <u>ilvG</u>, <u>ilvM</u>, <u>ilvE</u>, <u>ilvD</u> and <u>ilvA</u> genes, which encodes a large subunit and a small subunit of isozyme II of acetohydroxy-acid synthase, transaminase, dihydroxy-acid dehydratase and threonine deaminase, respectively. Of these enzymes, acetohydroxy-acid synthase, transaminase and dihydroxy-acid dehydratase catalyze the synthetic pathways from pyruvic acid to L-valine and from 2-ketobutyric acid to L-isoleucine, and threonine deaminase catalyzes the deamination from L-threonine to 2-ketobutyric acid, which is a rate-limiting step of L-isoleucine biosynthesis. Therefore, to conduct the reaction of L-valine synthesis so that it proceeds efficiently, an operon which does not express active threonine deaminase is used preferably. As such <u>ilvGMEDA</u> operons, an <u>ilvGMEDA</u> operon in which a mutation leading to the production of an inactivated threonine deaminase is introduced to <u>ilvA</u> or <u>ilvA</u> is destroyed, or an <u>ilvGMED</u> operon in which <u>ilvA</u> is deleted, can be used.

Because of the regulation of the expression (attenuation) of an <u>ilvGMEDA</u> operon by L-valine and/or L-isoleucine and/or L-leucine, the region necessary for the attenuation is preferably deleted or mutated to release the regulation of the expression caused by the generating L-valine.

The aforementioned <u>ilvGMEDA</u> operon, which does not express threonine deaminase activity and whose attenuation is released, can be obtained by mutating a wild-type <u>ilvGMEDA</u> operon or modifying it with genetic recombination techniques

The <u>ilvGMEDA</u> operon can include an operon derived from a microorganism belonging to the genus <u>Escherichia</u>, and particularly an <u>ilvGMEDA</u> operon derived from <u>E. coli</u>. Among microorganisms belonging to the genus <u>Escherichia</u>, the microorganism to be used is not particularly limited, however specifically, microorganisms described by Neidhardt, F.C. et al. (<u>Escherichia coli</u> and <u>Salmonella typhimurium</u>, American Society for Microbiology, Washington D.C., 1208, Table 1) can be utilized. When a wild-type strain is used as the donor strain of DNA including an <u>ilvGMEDA</u> operon, DNA including a wild-type <u>ilvGMEDA</u> operon is obtained.

However, when <u>E. coli</u> is used as the <u>DNA</u> donor strain of a wild-type <u>ilvGMEDA</u> operon, a wild-type K-12 strain does not express active isozyme II of acetohydroxy acid synthase (AHASII) because an <u>ilvG</u> gene possesses a frameshift mutation (Proc. Natl. Acad. Sci. USA, <u>78</u>, 922, 1991). Therefore, when the K-12 strain is used as the DNA donor strain, it is necessary that a mutant, in which the frame is restored so as to recover the activity of the <u>ilvG</u> geneencoded acetohydroxy-acid synthase, is prepared, and then used as the DNA donor strain. Otherwise, by using <u>E. coli</u>, other than a strain derived from the K-12 strain, as a DNA donor, only the <u>livG</u> gene may be isolated and introduced in <u>ilvGMEDA</u> operon derived from the K-12 strain. Thus, the <u>ilvMEDA</u> region is isolated from the K-12 strain as a DNA

donor, only the <u>ilvG</u> gene is isolated from <u>E. coli</u>, other than a strain derived from the K-12 strain, as a DNA donor, and the obtained both sequences are ligated together to form the full-length <u>ilvGMEDA</u> operon. The isozyme II of acetohydroxy-acid synthase (AHASII) is composed of two different large and small subunits. The large subunit is encoded by a <u>ilvG</u> gene. The small subunit is encoded by a <u>ilvM</u> gene.

The method of obtaining the <u>ilvGMEDA</u> operon which is released from the attenuation may be as follows:

The localization and DNA sequences of the attenuator which is 5'-upstream in the <u>ilvGMEDA</u> operon have been reported by R.P. Lawther et al. (Nucleic Acids Res., <u>15</u>, 2137 (1987)).

Starting from <u>ilvGMEDA</u> which does not express active threonine deaminase, an <u>ilvGMEDA</u> operon wherein active threonine deaminase is not expressed and attenuator is deleted is obtained by preparing an <u>ilvGMEDA</u> operon wherein the attenuator is deleted.

The nucleotide sequence shown in SEQ ID NO:1 is a sequence including the promoter, the attenuator and the <u>ilvG</u> gene-coding region among nucleotide sequences of the <u>ilvGMEDA</u> operon and includes a region necessary for attenuation. An amino acid sequence coded by the <u>ilvG</u> gene is shown in SEQ ID NO:2. Nucleotides 966 to 971 of the DNA sequence encode two consecutive leucine residues localized in the leader peptide, nucleotides 999 to 1007 of the DNA sequence encode three consecutive valine residues localized in the leader peptide, and nucleotides 1008 to 1016 of the DNA sequence encode three consecutive isoleucine residues localized in the leader peptide. Nucleotides 1081 to 1104 of the DNA sequence encode a portion forming a rho-independent terminator-like stem-loop structure in the attenuator.

Sufficient amounts of L-isoleucine, L-valine and L-leucine in a cell lead to the formation of the rho-independent terminator-like stem-loop structure with the RNA which is a transcript encoded by nucleotides 1081 to 1104 of the DNA sequence so that RNA polymerase terminates transcription, which represses the expression of the <a href="https://liven.com/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven

For example, the shortage of L-valine in a cell results in the shortage of L-valine-binding tRNA, which causes ribosomal translation to be retarded at consecutive valine residues localized in the region encoding the leader peptide. This leads to the formation of an additional mRNA configuration in the three-dimensional structure, resulting in the formation of rho-independent terminator-like stem-loop structure in the RNA, which is stimulated by the transcription of nucleotides 1081 to 1104 the DNA sequence, is repressed. Thus, RNA polymerase continues transcription, which results in the expression of the <a href="https://liven.com/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liven/liv

Therefore, to delete the region necessary for attenuation by L-valine, nucleotides 999 to 1007 or 1081 to 1104 of the DNA sequence shown in SEQ ID NO:1 may be deleted. Similarly, to delete the region necessary for attenuation by L-leucine in the production of L-leucine-producing microorganism as described below, nucleotides 966 to 971 or 1081 to 1104 of the DNA sequence disclosed in SEQ ID NO:1 may be deleted.

Thus, concepts of this deletion of the region necessary for attenuation include the insertion of an additional DNA fragment into the attenuator as well as the deletion of all parts or vicinities of attenuators which are 5'-upstream in the <a href="https://links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/links.com/

## (i) Isolation of wild-type ilvGMEDA operon

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To isolate a DNA containing the <u>ilvGMEDA</u> operon, the method, wherein <u>ilvGM</u>, <u>ilvE</u>, <u>ilvD</u> and <u>ilvA</u> genes are each isolated and then ligated, may be suggested. However, in the construction of an L-valine-producing microorganism, an <u>ilvA</u> gene encoding threonine deaminase is not necessary, so that <u>ilvGM</u>, <u>ilvE</u> and <u>ilvD</u> genes may be ligated to obtain a DNA including the <u>ilvGMED</u>.

First, E. coli, e.g. E. coli K-12, E. coli W3110, E. coli MC1061 (all of which include a frameshifted iivG), E. coli MI162 (thr-10, car-94, λ-, relA1, iivG603, thi-1) or E. coli B (the latter two of which include a normal iivG), is cultured to obtain the cultured cells. The microorganism may be cultured by the usual solid medium method, and preferably cultured according to the liquid medium method in consideration of efficiency in cell harvesting. A medium wherein yeast extracts, pepton, trypton or meat extracts are added to sodium chloride (NaCl) is to be used. Specifically, L-broth (Bacto-trypton 1%, Bacto-yeast extracts 0.5%, NaCl 0.5%, glucose 0.1%, pH 7.2) is to be used. The initial pH of the medium is preferably adjusted to 6-8. The cultivation is conducted at 30 to 42°C, preferably about 37°C, for 4-24 hours, with aeration, stirring and submerged in culture, with a shaking culture, or with a stationary culture. E. coli MI162 is available from the E. coli Genetic Stock Center (Connecticut, U.S.A.). The ID No. of this strain is CGSC5919. The detail

characteristics of this strain are described in Mol. Gen. Genet., 143, 243 (1976), and J. Bacteriol., 149, 294 (1982).

Thus, the obtained culture is centrifuged, e.g. at 3,000 r.p.m. for 5 minutes to obtain a pellet of <u>E. coli</u>. From this pellet, chromosomal DNA can be obtained by the method of Saitoh and Miura (Biochem. Biophys. Acta., <u>72</u>, 619 (1963)) or K. S. Kirby (Biochem. J., 64, 405 (1956)).

To isolate the <u>ilvGMEDA</u> operon from the resulting chromosomal DNA, the chromosomal DNA library is prepared. First, the chromosomal DNA is partially digested by a proper restriction enzyme to obtain a mixture of different DNA fragments. A wide variety of restriction enzymes can be used if the digestion reaction is adjusted for the degree of digestion. For example, the chromosomal DNA is digested with <u>Sau</u>3Al at not less than 30°C, preferably at 37°C, at an enzyme concentration of 1-10 units/ml for a varying period of time (1 minute to 2 hours).

Subsequently, the digested DNA was ligated to vector DNA which allows autonomous replication to prepare recombinant DNA. Specifically, the vector DNA is completely digested and cleaved with the restriction enzyme, e.g. <u>Bam</u>HI, which generates the restriction termini identical to those generated by Sau3Al used in the digestion of the chromosomal DNA, at a temperature of above 30°C, at a enzyme concentration of 1-100 units/ml for above 1 hour, preferably 1-3 hours. And then, the chromosomal DNA fragments and the cleaved vector DNA, obtained as mentioned above, were mixed, added to DNA ligase, preferably T4 DNA ligase, and reacted at a temperature of 4-16°C, at a enzyme concentration of 1-100 units/ml above 1 hour, preferably 6-24 hours to obtain recombinant DNA.

Using the resulting recombinant DNA, a microorganism belonging to the genus Escherichia, for example, a mutant which is deficient in acetohydroxy acid synthase activity such as E. coli Ml262 (leuB6, ilvI614, ilvH612,  $\lambda^2$ , relA1, spoT1, ilvB619, ilvG603, ilvG605(am), thi-1), transaminase B-deficient mutants such as E. coli AB2070 (proA2, trp-3, higG4, ilvE12, metE12, metE46, thi-1, ara-9, lac-Y1 or lacZ4, galK2, malA1, mtl-1, rpsL8 or rpsL9, ton-1, tsx-3,  $\lambda^R$ ,  $\lambda^2$ , supE44), or dihydroxy acid dehydratase-deficient mutants such as E. coli AB1280 (hisG1, ilvD16, metB1, argH1, thi-1, ara-13, lacY1 or lacZ4, gal-6, xyl-7, mtl-2, malA1, repsL8, 9, or 17, tonA2,  $\lambda^R$ ,  $\lambda^2$ , supE44), is transformed to prepare chromosomal DNA library. This transformation can be performed by the method of D.M. Morrison (Methods in Enzymology 68, 326, 1979) or a method wherein the treatment of a recipient cell with calcium chloride increases the permeability of DNA (Mandel, M. and Higa, A., J. Mol., Biol., 53, 159 (1970). E. coli Ml262 is available from the E. coli Genetic Stock Center (Connecticut, U.S.A.). The ID No. of this strain is CGSC5769. The detailed characteristics of this strain are described in Mol. Gen. Genet., 156, 1 (1977). E. coli AB2070 is available from the E. coli Genetic Stock Center (Connecticut, U.S.A.). The ID No. of this strain is CGSC2070. The detailed characteristics of this strain are described in J. Bacteriol., 109, 730 (1972).

Because the nucleotide sequence of the full-length <u>ilvGMEDA</u> operon has been reported (Nucleic Acids Res., <u>15</u>, 2137 (1987)), a certain length of DNA fragments including the aimed gene can be prepared by digesting the chromosomal DNA with a specific restriction enzyme. Only DNA fragments of a certain length are ligated to the vector DNA to generate recombinant DNA and to prepare the chromosomal DNA library, whereby the DNA fragment including the aimed gene can be obtained more efficiently.

From the obtained chromosomal DNA library, the strain which has recombinant DNA including the <a href="itvGM">itvGM</a> gene is obtained by selecting a strain having increased acetohydroxy acid synthase activity or a strain whose nutrient requirement caused by the deficient in acetohydroxy acid synthase gene is complemented.

From the obtained chromosomal DNA library, the strain which has recombinant DNA including an <a href="https://library.com/library">ilvE</a> gene is obtained by selecting a strain having increased transaminase B activity or a strain whose nutrient requirement caused by the deficiency in the transaminase B gene is complemented.

From the obtained chromosome DNA library, the strain which has recombinant DNA including an <a href="mailto:ilvD">ilvD</a> gene is obtained by selecting a strain having increased dihydroxy acid dehydratase activity or a strain whose nutrient requirement caused by the deficiency in the dihydroxy acid dehydratase gene is complemented.

To examine whether candidates to have recombinant DNA including the <u>ilvGM</u> gene have recombinant DNA wherein <u>ilvGM</u> gene is cloned or not, increase of acetohydroxy acid synthase activity is confirmed by preparing a cell extract from the candidate and further preparing a crude enzyme solution from this extract. The assay of acetohydroxy acid synthase activity can be performed by the method of M.D. Felice et al. (Methods in Enzymology <u>166</u>, 241).

Because the AHAS-deficient strain exhibits isoleucine, leucine and valine requirements, when the acetohydroxy acid synthase-deficient mutant is used as a host cell, DNA fragments including the <a href="https://live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/live.com/

Otherwise, DNA sequence containing the <u>ilvGM</u> gene has been reported by R.P. Lawther et al. (Nucleic Acids Res., <u>15</u>, 2137 (1987)). Thus, the confirmation can be performed by isolating the recombinant DNA from the candidates, by sequencing and comparing it with that described in the report.

As described above, there is a mutation within the open reading frame of the <u>ilvG</u> gene of <u>E. coli</u> K-12. As a result, the generated frameshift and further the emergence of a termination codon cause translational termination. Thus, the termination codon emerges at a position of 982-984 downstream of the initiation codon ATG (at a position of 1-3) of the <u>ilvG</u> gene. Therefore, when the <u>ilvGM</u> gene obtained from the strain is used, the mutation region needs to back to the normal sequence by the site-directed mutagenesis method. For example, for the <u>ilvG</u> gene (<u>ilvG603</u>) of <u>Escherichia coli</u>

MI162, the frame is normalized by placing two base pairs of TG before the termination codon TGA at a position of 982-984. The other mutations are described in Fig.2 in J. Bacteriol., 149, 294 (1982).

The method for confirming whether candidates to have recombinant DNA including the <u>ilvE</u> gene have recombinant DNA wherein <u>ilvE</u> gene is cloned or not is as follows. Because the transaminase B-deficient mutant exhibits an isoleucine requirement, when the transaminase B-deficient mutant is used as a host cell, DNA fragments including the <u>ilvE</u> gene can be obtained by isolating a strain which can grow in a minimum medium without isoleucine, and by collecting recombinant DNA from said strain.

Otherwise, the DNA sequence including the <u>ilvE</u> gene has been reported by R.P. Lawther et al. (Nucleic Acids Res., <u>15</u>, 2137 (1987)). Thus, the confirmation can be performed by isolating the recombinant DNA from the candidates, by sequencing and comparing it with that described in the report.

The method for confirming whether candidates to have recombinant DNA including the <u>ilvD</u> gene have recombinant DNA wherein <u>ilvD</u> gene is cloned or not is as follows. Because the dihydroxy acid dehydratase-deficient mutant exhibits an isoleucine, leucine and valine requirement, when the dihydroxy acid dehydratase-deficient mutant is used as a host cell, DNA fragments including the <u>ilvD</u> gene can be obtained by isolating a strain which can grow in a minimum medium without valine, and by collecting recombinant DNA from said strain.

Otherwise, the DNA sequence including the <u>ilvD</u> gene has been reported by R.P. Lawther et al. (Nucleic Acids Res., 15, 2137 (1987)). Thus, the confirmation can be performed by isolating the recombinant DNA from the candidates, by sequencing and comparing it with that described in the report.

From each of the aforementioned strains, recombinant DNA can be isolated e.g. by the methods of P. Guerry et al. (J. Bacteriol., <u>166</u>, 1064 (1973)) and D.B. Clewell (J. Bacteriol., <u>110</u>, 667 (1972)).

To obtain a full-length <u>ilvGMEDA</u> operon, a DNA fragment including the <u>ilvGM</u> gene, a DNA fragment including the <u>ilvE</u> gene, and a DNA fragment including the <u>ilvD</u> gene are ligated. In the ligation, the DNA sequence of the full-length <u>ilvGMEDA</u> described by R.P. Lawther (Nucleic Acids Res., <u>15</u>, 2137 (1987)) can be used as a reference.

A wild-type <u>ilvGMEDA</u> operon may be obtained by preparing chromosomal DNA from a strain having the wild-type <u>ilvGMEDA</u> in its chromosome by the method of Saitoh and Miura, and by amplifying the <u>ilvGMEDA</u> operon by the polyrmrase chain reaction method (PCR; see White, T.J. et al.; Trends Gent., <u>5</u>, 185 (1989)). As a DNA primer in amplification, those complementary to both 3'-ends of DNA double strands including all or a part of the region of <u>ilvGMEDA</u> operon are used. In the amplification of only a part of the region of the <u>ilvGMEDA</u> operon, DNA fragments including the entire region are screened by using said DNA fragment as a probe. In the amplification of the entire region of the <u>ilvGMEDA</u> operon, an agarose gel electrophoresis of a PCR solution which contains DNA fragments including the amplified <u>ilvGMEDA</u> operon followed by extraction of the aimed DNA fragments allows for the collection of DNA fragments including the <u>ilvGMEDA</u> operon. Because in this case, also, the <u>ilvA</u> gene is not essential to the construction of a L-valine-producing microorganism, only the <u>ilvGMEDA</u> may be amplified.

When preparing a DNA primer, the DNA sequence of the full-length <u>ilvGMEDA</u> operon described by R.P. Lawther et al. (Nucleic Acids Res., <u>15</u>, 2137 (1987)) can be used as a reference.

Primer DNA can be synthesized in a commercially available DNA synthesizer (e.g. Applied Biosystems, DNA synthesizer model 380B) by the phosphoramidite method (Tetrahedron Letters, <u>22</u>, 1859 (1981)). PCR can be performed in a commercially available PCR system (Perkin Elmer, DNA thermal cycler PJ2000), using Taq DNA polymerase (supplied by Takara Shuzo, Ltd.) according to the method indicated by suppliers.

The <u>ilvGMEDA</u> operon amplified by the PCR method is ligated to vector DNA, which allows autonomous replication, in the cell of the microorganism belonging to the genus <u>Escherichia</u> and induced into the cell of a microorganism belonging to the genus <u>Escherichia</u>, whereby the induction of a mutation into the <u>ilvA</u> gene and the deletion of the region necessary for attenuation are facilitated. The vector DNA, transformation method, and further confirmation of the <u>ilvGMEDA</u> operon are the same as described above.

When <u>E. coli</u> K-12, <u>E. coli</u> W3110 and <u>E. coli</u> MC1061 are used as the donor microorganism of the <u>ilvGMEDA</u> operon, because a frameshift mutation is present within the open reading frame of the <u>ilvG</u> gene, this mutation needs to be normalized by the site-directed mutagenesis method. When <u>E. coli</u> MI162 (thr-10, car-94,  $\lambda^2$ , relA1, ilvG603, thi-1), and <u>E. coli</u> B are used as the donor microorganism of the <u>ilvGMEDA</u> operon, the <u>ilvG</u> gene can be used as it is.

# (ii) Deletion of the region of ilvGMEDA operon necessary for attenuation

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Concepts of the deletion of the region necessary for attenuation from <u>ilvGMEDA</u> include an insertion of an additional DNA fragment into the attenuator as well as a deletion of all of, a part of or an area surrounding the attenuator which is upstream in the <u>ilvGMEDA</u> operon. Herein, the "attenuator" means a DNA sequence which forms a rho-independent terminator-like stem-loop structure. For example, the sequence corresponds to nucleotides 1081 to 1104 of the DNA sequence shown in SEQ ID NO:1.

To delete the attenuator, DNA fragments upstream and downstream of the attenuator in the <u>ilvGMEDA</u> operon may be each prepared to ligate the both DNA fragments. For example, the DNA fragment upstream of the attenuator in the

<u>ilvGMEDA</u> operon can be prepared by cleaving a DNA fragment including the full length of the <u>ilvGMEDA</u> operon with a proper restriction enzyme. Otherwise, the DNA fragment upstream of the attenuator in the <u>ilvGMEDA</u> operon may be amplified by the PCR method. The primer DNA used in the PCR method may be chemically synthesized on the basis of the DNA sequences described by R.P. Lawther et al. (Nucleic Acids Res., <u>15</u>, 2137 (1987)) and G.Coopola et al. (Gene, <u>97</u>, 21 (1991)). Furthermore, the DNA fragment upstream of the attenuator in the <u>ilvGMEDA</u> operon may be chemically synthesized.

The method for preparing the DNA fragment downstream of the attenuator in the <u>ilvGMEDA</u> operon is similar to that above.

Starting from the <u>ilvGMEDA</u> operon, the <u>ilvGMEDA</u> operon wherein part or vicinity of the attenuator is deleted may be prepared. Because the location and DNA sequence of the attenuator have been reported by R.P. Lawther et al. (Nucleic Acids Res., <u>15</u>, 2137 (1987)), DNA to be deleted is determined on the basis of the sequence.

The DNA to be deleted is preferably the DNA sequence which is necessary to form a rho-independent terminator-like stem-loop structure, and/or includes the region encoding consecutive valine residues which are upstream in the stem-loop structure. To delete a part of or an area around the attenuator, DNA fragments upstream and downstream of the attenuator in the <a href="ilvGMEDA">ilvGMEDA</a> operon may be each prepared to ligate both DNA fragments. For example, the DNA fragment upstream in the attenuator of the <a href="ilvGMEDA">ilvGMEDA</a> operon can be prepared by cleaving a DNA fragment including the full length of the <a href="ilvGMEDA">ilvGMEDA</a> operon with a proper restriction enzyme. Otherwise, the DNA fragment upstream of the DNA to be deleted in the <a href="ilvGMEDA">ilvGMEDA</a> operon may be amplified by the PCR method. The primer DNA used in the PCR method may be chemically synthesized on the basis of the DNA sequences described by R.P. Lawther et al. (Nucleic Acids Res., <a href="15">15</a>, 2137 (1987)) and G. Coopola et al. (Gene, <a href="15">97</a>, 21 (1991)). Furthermore, the DNA fragment upstream of the DNA to be deleted in the <a href="ilvGMEDA">ilvGMEDA</a> operon may be chemically synthesized.

The method for preparing the DNA fragment downstream of the DNA to be deleted in the <u>ilvGMEDA</u> operon is similar to that above.

Starting from the <u>ilvGMEDA</u> operon, an <u>ilvGMEDA</u> operon wherein an additional DNA fragment is inserted into the attenuator may be prepared. Because the location and DNA sequence of the attenuator have been reported by R.P. Lawther et al. or G. Coppola et al., the position of the insertion and the DNA sequence of an additional DNA fragment to be inserted are determined on the basis of the sequence.

The additional DNA fragment to be inserted is preferably inserted into the DNA sequence which is necessary to form a rho-independent terminator-like stem-loop structure, or into the DNA region encoding consecutive valine residues which is upstream of the stem-loop structure. As a result of the insertion, the attenuator can not form a rho-independent terminator-like stem-loop structure and so the attenuator is expected to loose its function.

The DNA sequence of the additional DNA fragment to be inserted is preferably designed not to form a rho-independent terminator-like stem-loop structure, and to cause the consecutive valine residues not to be present upstream of the rho-independent terminator-like stem-loop structure when inserted.

To insert an additional DNA fragment into the attenuator, the DNA fragment of the <a href="ilvGEMED">ilvGEMED</a> operon which is upstream of the additional DNA fragment to be inserted, the DNA fragment of the <a href="ilvGEMED">ilvGEMED</a> operon which is downstream of the additional DNA fragment to be inserted may be prepared to ligate these three DNA fragments. For example, the DNA fragment upstream of the additional DNA fragment in the <a href="ilvGMEDA">ilvGMEDA</a> operon can be prepared by cleaving a DNA fragment including the full length of the <a href="ilvGMEDA">ilvGMEDA</a> operon with a proper restriction enzyme. Otherwise, the DNA fragment of the <a href="ilvGMEDA">ilvGMEDA</a> operon which is upstream of the additional DNA fragment may be amplified by the PCR method. The primer DNA used in the PCR method may be chemically synthesized on the basis of the DNA sequences described by R.P. Lawther et al. (Nucleic Acids Res., <a href="15">15</a>, 2137 (1987)) and G. Coopola et al. (Gene, <a href="97">97</a>, 21 (1991)). Furthermore, the DNA fragment upstream of the additional DNA fragment in the <a href="ilvGMEDA">ilvGMEDA</a> operon may be chemically synthesized.

The method for preparing the DNA fragment downstream of the additional DNA fragment in the <u>ilvGMEDA</u> operon is similar to that above.

The additional DNA fragment to be inserted can be prepared by chemical synthesis.

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In the amplification of the DNA fragment of the <u>ilvGMEDA</u> operon which is upstream of the DNA region into which the additional DNA fragment, or the DNA fragment of the <u>ilvGMEDA</u> operon which is downstream of the DNA region into which the additional DNA fragment, the additional DNA fragment to be inserted can be ligated with the primer DNA. For example, the 3'-end DNA primer used for the amplification of the DNA fragment upstream of the DNA region into which the additional DNA fragment is ligated with one of the strands of the additional DNA fragment to be inserted. Similarly, the 5'-end DNA primer used for the amplification of the DNA fragment downstream of the DNA region into which the additional DNA fragment is ligated with the complementary one of the strands of the additional DNA fragment to be inserted. Two different DNA fragments which have been amplified using above-mentioned primers are ligated.

### (iii) Inactivation of threonine deaminase

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When the obtained <u>ilv</u> operon contains an <u>ilvA</u> gene, the <u>ilvA</u> is deleted, or modified to cause a mutation, insertion and deletion within the <u>ilvA</u> so as to inactivate the expressed threonine deaminase. As a modification, for example, a restriction site in the <u>ilvA</u> gene can be cleaved to delete a DNA fragment which is downstream in the cleaved site. A DNA fragment may be cut out by cleaving the <u>ilvA</u> gene at two sites and then re-ligating it. Further, the expressed threonine deaminase can be inactivated by inserting another DNA fragment such as a synthesized DNA into the restriction site. When the restriction sites are cohesive ends, these cohesive ends are treated so as to be blunt ends, and then the resulting ends are ligated together, whereby the expressed threonine deaminase can be inactivated. Furthermore, the expressed threonine deaminase can be inactivated by site-specific mutagenesis and the like.

Hereinafter, the <u>ilvGMEDA</u> operon, wherein attenuation is repressed and the threonine deaminase activity is not expressed, or <u>ilvA</u> is deleted, is referred to as a derepressed <u>ilvGMEDA\*</u> operon, where A\* represents the deleted <u>ilvA</u> gene, or the <u>ilvA</u> encoding inactivated threonine deaminase or a part thereof.

## (iv) Introduction of derepressed ilvGMEDA\* operon in a microorganism belonging to the genus Escherichia

A DNA fragment including the derepressed <u>ilvGMEDA\*</u> operon obtained as above used as a recombinant DNA, introduced into a proper host microorganism and expressed, whereby the microorganism, in which the expression of the enzymes involving valine biosynthesis encoded by the <u>ilvGMEDA\*</u> operon is enhanced, can be obtained. As a host microorganism, a microorganism belonging to the genus <u>Escherichia</u>, e.g. <u>Escherichia coli</u>, is preferably used.

A derepressed <u>ilvGMEDA</u>\* operon, which is cut out from a recombinant DNA and inserted into the other vector DNA, may be used. As the vector DNA which can be used in the present invention, for example, pUC19, pUC18, BR322, pHSG299, pHSG399, pHSG399, pHSG399, pHSG398, RSF1010, pMW119, pMW118, pMW219 and pMW218 can be used. Additionally, a vector of phage DNA also can be used.

Further, to efficiently perform the expression of the derepressed <u>ilvGMEDA</u>\* operon, the other promoters, which act in a microorganism including lac, trp and  $P_L$ , may be ligated, and the promoter innate to the <u>ilvGMEDA</u>\* operon may be used as it is or after amplification.

As mentioned above, the DNA fragment including the derepressed <u>ilvGMEDA</u>\* operon may be present in a host microorganism as an extrachromosomal DNA such as a plasmid by inserting the operon into vector DNA which allows autonomous replication into the host, while the derepressed <u>ilvGMEDA</u>\* operon may be inserted into the chromosome of the host microorganism by using the techniques of transduction, a transposon (Berg, D.E. and Berg, C. M., Bio/Technol., <u>1</u>, 417 (1983)), a Mu phage (Japanese Patent Application Laid-Open No. 2-109985(1990)) or by homologous recombination (Experiments in Molecular Genetics, Cold Spring Habor Lab. (1972)). The number of the derepressed <u>ilvGMEDA</u>\* operons introduced into the host may be either one or more.

As described above, an L-valine producing microorganism can be obtained by introducing the DNA fragment including the derepressed <u>ilvGMEDA\*</u> operon into a lipoic acid requiring and/or H\*-ATPase-deficient microorganism belonging to the genus <u>Escherichia</u>. Also, an L-valine-producing microorganism can be obtained by introducing lipoic acid requirement and/or H\*-ATPase deficiency to a microoranism belonging to the genus <u>Escherichia</u> and carrying the DNA fragment including the derepressed <u>ilvGMEDA\*</u> operon.

#### (2) L-leucine-producing microorganism

As illustrated in the example below, it is found that a lipoic acid-requiring or H<sup>+</sup>-ATPase-deficient microorganism belonging to the genus <u>Escherichia</u> can increase the L-valine productivity. This finding suggests that a lipoic acid-requiring mutation or H<sup>+</sup>-ATPase-deficient mutation causes the intracellular metabolism to stimulate the L-valine synthesis. Therefore, the lipoic acid requiring mutation or H<sup>+</sup>-ATPase-deficient mutation is considered to promote L-leucine biosynthesis whose synthetic pathway branches out from the final intermediate of L-valine. Thus, if the capability of producing L-leucine is added to or enhanced in a lipoic acid-requiring mutant, a H<sup>+</sup>-ATPase-deficient mutant, or a lipoic acid-requiring and H<sup>+</sup>-ATPase-deficient mutant, the capability of producing L-leucine is expected to be added thereto or enhanced.

The addition or enhancement of the capability of producing L-leucine, for example, is performed by introducing an L-leucine biosynthetic gene, wherein the regulatory mechanism is substantially released, into a microorganism belonging to the genus <u>Escherichia</u>, in addition to the properties necessary for the production of L-valine. And a mutation, whereby the regulatory mechanism of L-leucine biosynthesis in a microorganism belonging to the genus <u>Escherichia</u> is substantially released, may be introduced. These genes can include, for example, a leuA gene in which inhibition by L-leucine is substantially released.

In addition to the aforementioned capability of producing L-valine or L-leucine, a microorganism of the present invention may have the known characteristics which are effective in enhancing its capability of producing an amino acid,

for example, various nutrient requirements, resistance to drugs, sensitivity to drugs, and drug dependence, or characteristics wherein a gene promoting the biosynthesis of an amino acid is amplified by means of gene engineering.

(2) Production of L-valine or L-leucine of the present invention

The production of L-valine or L-leucine of the invention can be performed by culturing the microorganism of the present invention in a liquid medium, to allow L-valine or L-leucine to be produced and accumulated in the liquid medium, and collecting L-valine or L-leucine from this liquid medium. In this production, the L-valine-producing microorganism of the present invention is used in the production of L-valine, and the L-leucine producing microorganism of the present invention is used in the production of L-leucine.

In the producing method of the present invention, the cultivation of the L-valine or L-leucine-producing microorganism, the collection and purification of L-valine or L-leucine from the liquid medium may be performed in a manner similar to the conventional fermentation method wherein an amino acid is produced using a microorganism. A medium used for culture may be either a synthetic medium or a natural medium, so long as the medium includes a carbon source and a nitrogen source and minerals and, if necessary, appropriate amounts of nutrients which the microorganism requires for growth. The carbon source may include various carbohydrates such as glucose and sucrose, and various organic acids. Depending on the mode of assimilation of the used microorganism, alcohol including ethanol and glycerol may be used. As the nitrogen source, various ammonium salts such as ammonia and ammonium sulfate, other nitrogen compounds such as amines, a natural nitrogen source such as peptone, soybean-hydrolysate and digested fermentative microorganism are used. As minerals, potassium monophosphate, magnesium sulfate, sodium chloride, ferrous sulfate, manganese sulfate, calcium carbonate, etc. are used.

The cultivation is performed preferably under aerobic conditions such as a shake culture, and an aeration and stirring culture, at a temperature of 20 to 40°C, preferably 30 to 38°C. The pH of the culture is usually between 5 and 9, preferably between 6.5 and 7.2. The pH of the culture can be adjusted with ammonia, calcium carbonate, various acids, various bases, and buffers. Usually, a 1 to 3-day cultivation leads to the accumulation of the target L-valine or L-leucine in the liquid medium.

After cultivation, solids such as cells can be removed from the liquid medium by centrifugation and membrane filtration, and then the target L-valine or L-leucine can be collected and purified by ion-exchange, concentration and crystallization methods.

## Brief Description of the Drawings

Figure 1 is a scheme of the construction of the plasmid pHSGSK.

Figure 2 is a scheme of the construction of the plasmid pdGM1.

Figure 3 is a scheme of the construction of the plasmid pMWGMA2.

Figure 4 is a scheme of the construction of the plasmid pMWD5.

Figure 5 is a scheme of the construction of the plasmid pMWdAR6.

## Best Mode fo Carrying Out the Invention

The invention is described with reference to the following example:

## Example 1

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- Creation of L-valine-producing microorganism
  - (1) Construction of pMWdAR6 carrying the derepressed ilvGMEDA\* operon

The chromosomal DNA was extracted from <u>Escherichia coli</u> MI162. The chromosomal DNA was cleaved with restriction enzyme <u>HindIII</u>. The length of a <u>HindIII-HindIII</u> DNA fragment including <u>ilvGM</u> genes was found to be 4.8 kb. Therefore, the <u>HindIII-HindIII</u> DNA fragment with approximately 4.8 kb and the DNA fragment obtained by digestion of the plasmid vector pBR322 (purchased form Takara Shuzo, Co., Ltd.) with <u>HindIII</u>, were ligated.

The resulting DNA-ligated mixture was induced into Escherichia coli MI162 which is an acetohydroxy-acid synthase-deficient strain. The strains in which the deficiency of acetohydroxy-acid synthase was complemented by transformation were selected and the plasmid structure was isolated from the selected strains. The results of the analysis of the plasmid revealed that a 4.8-kb DNA fragment containing the <u>ilvGM</u> gene and a portion of 5'-terminal of <u>ilvE</u> gene was inserted into the <u>HindIII</u> site of the pBR322. The plasmid was termed pBRGM7.

The synthetic oligonucleotides shown in SEQ ID NO:3 and NO:4 were synthesized with reference to the DNA

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sequence of the <u>ilvGM</u> gene described in Gene, <u>97</u>, 21, (1991), Pro. Natl. Acad. Sci. U.S.A., <u>78</u>, 922, (1981) and J. Bacteriol., <u>149</u>, 294, (1982). DNA was amplified by the PCR method, using both synthesized DNAs as a primer and chromosomal DNA of MI162 strain as a template. The amplified DNA fragment included nucleotides 25 to 952 of the nucleotide sequence shown in SEQ ID NO:1. The fragment was termed Fragment (A).

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Similarly, the synthetic oligonucleotides shown in SEQ ID NO:5 and NO:6 were synthesized with reference to the DNA sequence described in Gene, <u>97</u>, 21, (1991), Pro. Natl. Acad. Sci. U.S.A., <u>78</u>, 922, (1981) and J. Bacteriol., <u>149</u>, 294, (1982). DNA was amplified by the PCR method, using both synthesized DNAs as a primer and chromosomal DNA of the MI162 strain as a template. The amplified DNA fragment included nucleotides 1161 to 2421 of the nucleotide sequence shown in SEQ ID NO:1. The fragment was termed Fragment (B).

The plasmid pUCA was prepared by ligating the large fragment obtained by digestion of Fragment (A) with <u>Smalland the DNA fragment obtained by digestion of the vector, pUC18 (Takara Shuzo, Co., Ltd.) with <u>Smalland the DNA fragment obtained by digestion of Fragment (B) with Kpnl and the DNA fragment obtained by digestion of the vector, pHSG399 (Takara Shuzo, Co., Ltd.) with <u>Hincll and Kpnl.</u></u></u>

The plasmid pUCA was digested with <u>KpnI</u>, the blunt-end fragment was prepared with the large fragment of DNA polymerase I (Klenow fragment), and digested with <u>PstI</u>, and finally, a DNA fragment containing Fragment (A) was isolated. Plasmid pHSGB was digested with HindIII, the blunt-end fragment was prepared with the large fragment of DNA polymerase I (Klenow fragment), and digested with <u>PstI</u>, and finally, a DNA fragment containing Fragment (B) was isolated. The plasmid pHSGSK was prepared by ligating both DNA fragments.

The <u>Smal-Kpn</u>I fragment derived from Fragments (A) and (B) in pHSGSK was termed Fragment (C). Fragment (C) corresponded to a fragment obtained by digestion of a 4.8-kb <u>HindIII-HindIII</u> fragment with <u>Smal</u> and <u>Kpn</u>I contained a promoter, the SD sequence and a upstream region of the <u>ilvG</u> gene, but lost the DNA sequence of 0.2 kb from a leader sequence to an attenuator. The scheme of construction of pHSGSK is summarized in Fig. 1.

Fragment (C) was obtained by digestion of the plasmid pHSGSK with <u>Smal</u> and <u>Kpnl</u>, the large DNA fragment was obtained by digestion of the plasmid pBRGM7 with <u>Smal</u> and <u>Kpnl</u>, and the both two fragments were ligated. The obtained plasmid was termed pdGM1. pdGM1 harbored a 4.6-kb <u>HindIII-HindIII</u> fragment including the <u>ilvGM</u> gene, which lost the region necessary for attenuation. This <u>ilvGM</u> gene which loses the region necessary for attenuation represents "AattGM". The scheme of the construction of pdGM1 is summarized in Figure 2.

The plasmid pDRIA4 described in Japanese Patent Application Laid-Open No. 2-458(1990) is prepared by combining the shuttle vector pDR1120, which allows autonomous replication in both a microorganism belonging to the genus Escherichia and a microorganism belonging to the genus Brevibacterium, with a BamHI-BamHI fragment including the ilvA gene encoding threonine deaminase and a portion of the 3'-terminal of the ilvD gene derived from E. coli K-12. Japanese Patent Application Laid-Open No. 2-458(1990) describes that the length of the BamHI-BamHI fragment is 2.3 kb; however, at present, the length of this fragment has been found to be 2.75 kb. The plasmid pDRIA4 is not present within the chromosomal DNA of Brevibacterium flavum AJ12358 (FERM P-9764) or Brevibacterium flavum AJ12359 (FERM P-9765). From these strains, the plasmid pDRIA4 can be prepared according to the usual method. The feedback inhibition of threonine deaminase encoded by the ilvA gene in pDRIA4 by L-isoleucine is released, whereas this release of the feedback inhibition is not essential in the present invention.

From a 2.75-kb <u>Bam</u>HI-<u>Bam</u>HI DNA fragment in the plasmid pDRIA4, a <u>Hin</u>dIII-<u>Bam</u>HI fragment including the <u>ilvA</u> gene encoding threonine deaminase, in which the inhibition by L-isoleucine was released, was prepared, and ligated to a DNA fragment obtained by cleaving the vector pMW119 (NIPPON GENE) with <u>Hin</u>dIII and <u>Bam</u>HI. The resulting plasmid was termed pMWA1.

A DNA fragment obtained by cleaving the plasmid pMWA1 with <u>HindIII</u> and a DNA fragment obtained by cleaving the plasmid pdGM1 with <u>HindIII</u> were ligated. According to the analysis of the position of the restriction sites of the ligated plasmids, the plasmid in which the transcriptional orientations of the <u>ilvGM</u> and <u>ilvA</u> genes were the same was selected, and termed pMWGMA2. The pMWGMA2 includes the <u>ilvGM</u> gene in which an attenuator was deleted, a 5'-terminal portion of the <u>ilvE</u> gene, and a 3'-terminal portion of the <u>ilvD</u> gene. The scheme of the construction of pMWGMA2 is summarized in Figure 3.

The chromosomal DNA of Escherichia coli MI162 was prepared and cleaved with Sall and Pstl to prepare the mixture of DNA fragments. On the other hand, a DNA fragment was prepared by cleaving the vector pUC19 (Takara Shuzo, Co., Ltd.) with Sall and Pstl. The mixture of DNA fragments was ligated to the DNA fragment obtained by cleaving pUC19, and the DNA mixture was obtained. The DNA mixture was induced into AB2070, a transaminase B-deficient strain, (provided from Escherichia coli Genetics Stock Center. J. Bacteriol., 109, 703, (1972), CGSC2070) and a transformant, in which the branched-chain amino-acid requirement was recovered, was selected. As a result of the preparation of a plasmid from the strain, the plasmid harbored a DNA fragment obtained by cleaving the plasmid pUC19 with Sall and Pstl, and a Sall-Pstl DNA fragment including the ilvE gene, which were ligated. The plasmid was termed pUCE1. The pUCE1 includes a 3'-terminal portion of the ilvM gene, the ilvE gene, and a 5'-terminal portion of the ilvD gene.

A DNA-fragment mixture was prepared by partially digesting pMWGMA2 with HindIII. On the other hand, a 1.7-kb

HindIII-HindIII DNA fragment containing a portion of the ilvE gene and a 5'-terminal portion of the ilvD gene was prepared by cleaving pUCE1 with HindIII. Using a DNA mixture obtained by ligating both of the DNA fragments, AB1280, a dihydroxy-acid dehydratase(ilvD gene product)-deficient strain, was transformed, and the strain which recovered branched chain amino acid requirement was selected from the transformants. In the plasmid prepared from the resulting transformant, a DNA fragment obtained by cleaving only the HindIII site between ΔattGM and ilvA of pMWGMA2 with HindIII, and a 1.7-kb HindIII-HindIII DNA fragment including a portion of the ilvE gene and a portion of the ilvD gene derived from pUCE1 were ligated, and the ilvGMEDA operon was reconstructed. The plasmid was termed pMWD5. The scheme of the construction of pMWD5 is summarized in Figure 4.

The resulting plasmid pMWD5 derived from the vector pMW119 harbors the <u>ilvGMEDA</u> operon in which the region necessary for attenuation is deleted.

Subsequently, the plasmid pMWD5 was completely digested with <u>SnaBl</u> and then partially digested with <u>Acc</u>III. The resulting DNA fragment was self-ligated to obtain the plasmid pMWdAR6 in which only the <u>ilvA</u> gene was destroyed (Figure 5). This plasmid pMWdAR6 includes the <u>ilvGMEDA</u> operon in which the region necessary for attenuation is deleted and the <u>ilvA</u> gene is destroyed.

### (2) Creation of L-valine-producing-microorganism

Using the plasmid pMWdAR6 carrying the <u>ilvGMED</u> operon which was obtained as described above, <u>E. coli</u> W1485lip2 (ATCC25645), a lipoic acid-requiring mutant; <u>E. coli</u> W1485atpA401, a H<sup>+</sup>-ATPase-deficient mutant; <u>E. coli</u> W1485 (ATCC12435) were each transformed and the following transformants were obtained:

- 1) E. coli W1485/pMWdAR6
- 2) E. coli W1485atpA401/pMWdAR6
- 3) E. coli W1485lip2/pMWdAR6
- 4) E. coli AJ12631/pMWdAR6

E. coli AJ12631 was obtained by transducing atpA401, a mutant gene, encoding mutant alpha subunit of F1 of H<sup>+</sup>-ATPase derived from E. coli AN718 (CGSC6308) into E. coli W1485lip2 (ATCC25645) (see Japanese Patent Application Laid-Open No. 5-137568(1993)). In selection of a transduced strain with a H<sup>+</sup>-ATPase-deficient mutation, bgl gene positioned in the vicinity of atpA401 gene was used as a marker. Since the bgl gene encodes phospho-beta-glucosidase, E. coli having the wild-type bgl gene (bgl<sup>+</sup>) cannot assimilate salicin, whereas E. coli having the mutant bgl gene (bgl<sup>+</sup>) can grow utilizing salicin as the sole carbon source, so that the colonies of a salicin-assimilating strain make a bromothymol blue-added medium plate turn yellow by an organic acid produced by the strain. Therefore, if the mutant bgl gene (bgl<sup>+</sup>) and atpA401 gene are linked-transduced, a H<sup>+</sup>-ATPase-deficient mutant can be selected efficiently. First, the salicin-assimilating (bgl<sup>+</sup>) strain was isolated from E. coli AN718, and then AN718 (bgl<sup>+</sup>) was infected by Plkc, and E. coli W1485lip2 was transduced using the obtained lysate. For the resulting transductant, a lipoic acid requirement and H<sup>+</sup>-ATPase activity was determined to confirm the presence of lipoic acid-requiring and H<sup>+</sup>-ATPase-deficient mutations.

Similarly, E. coli W1485atpA401 was obtained by transducing atpA401 into E. coli W1485.

## Example 2

### Production of L-valine

The L-valine productivity of L-valine-producing microorganism obtained in Example 1 was evaluated. Each of transformants was plated on the medium comprising Bacto-typtone 1%, yeast extract 0.5%, NaCl 0.5%, agar 1.5%, and ampicillin  $100~\mu g/ml$ , cultured at 37°C for 18 to 24 hours, and then a part of them was transferred to 20 ml of a fermentation medium (glucose 4%, ammonium sulfate 1.6%, potassium dihydrogen-phosphate 0.1%, magnesium sulfate heptahydrate 0.1%, ferrous sulfate heptahydrate 0.001%, manganese sulfate pentahydrate 0.001%, yeast extract 0.2%, Bacto-tryptone 0.2%, calcium carbonate 3%, pH 7.0) with a platinum transfer loop, and incubated at 37°C for 24 hours. In culture of lipoic acid-requiring mutant, lipoic acid was added at a final concentration of 1  $\mu g/L$ .

The concentration of L-valine in the supernatant of the culture, from which the cells were removed, was determined by high-performance liquid chromatography using a cation exchange column (CPK08: Asahi Chemical Industry Co., Ltd.). The results are shown in Table 1.

Table 1

Productivity of the L-vi	aline of each strain
E. coli transformant	Productivity of L-valine (g/L)
W1485	0.1
W1485/pMWdAR6	6.9
W1485atpA401/pMWdAR6	8.0
W1485lip2/pMWdAR6	7.8
AJ12631/pMWdAR6	9.2

The results reveal that when a DNA fragment including the <u>ilvDMEDA</u>\* operon in which threonine deaminase activity is not expressed and the region necessary for attenuation is deleted is introduced into a lipoic acid-requiring and/or H<sup>+</sup>-ATPase-deficient <u>E. coli</u> as a host cell, the resulting <u>E. coli</u> showed enhanced productivity of L-valine. If a lipoic acid-requiring and H<sup>+</sup>-ATPase-deficient strain is used as a host, the productivity of L-valine can be further enhanced.

Industrial Applicability

According to the present invention, it becomes possible to enhance the capability of L-valine or L-leucine production of a L-valine or L-leucine-producing microorganism. By using a microorganism of the present invention, L-valine and L-leucine can be produced efficiently.

## Sequence Listing

- (1) GENERAL INFORMATION:
  - (i) APPLICANT: AJINOMOTO CO., LTD.
  - (ii) TITLE OF INVENTION: METHODS FOR PRODUCING L-VALINE AND L-LEUCINE
  - (iii) NUMBER OF SEQUENCES: 6
  - (iv) CORRESPONDENCE ADDRESS:
    - (A) ADDRESSEE: Ajinomoto Co., Ltd.
    - (B) STREET: 15-1, Kyobashi 1-chome, Chuo-ku
    - (C) CITY: Tokyo 104
    - (D) STATE:
    - (E) COUNTRY: Japan
    - (F) ZIP:
  - (v) COMPUTER READABLE FORM:
    - (A) MEDIUM TYPE: Floppy disk
    - (B) COMPUTER: IBM PC compatible
    - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
    - (D) SOFTWARE: FastSEQ Version 1.5
  - (vi) CURRENT APPLICATION DATA:
    - (A) APPLICATION NUMBER:
    - (B) FILING DATE: 30.08.95
    - (C) CLASSIFICATION:
  - (viii) ATTORNEY/AGENT INFORMATION:
    - (A) NAME: Strehl Schübel-Hopf Groening & Partner
    - (B) REGISTRATION NUMBER: 94
    - (C) REFERENCE/DOCKET NUMBER: EPA-43613
    - (ix) TELECOMMUNICATION INFORMATION:
      - (A) TELEPHONE: [49](89)22-39 11
      - (B) TELEFAX: [49](89)22 39 15
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 2841 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(ii) MOLECULAR TYPE: genomic DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(vi) ORIGINAL SOURCE:	
(A) ORGANISM: Escherichia coli	
(B) STRAIN: MI162	
(ix) FEATURE:	
(A) NAME/KEY: CDS	
(B) LOCATION: 9571055	
(C) IDENTIFICATION METHOD: S	
(ix) FEATURE:	
(A) NAME/KEY: attenuator	
(B) LOCATION: 10811104	
(C) IDENTIFICATION METHOD: S	
(ix) FEATURE:	
(A) NAME/KEY: CDS	
(B) LOCATION: 11952841	
(C) IDENTIFICATION METHOD: S	
(ix) FEATURE:	
(A) NAME/KEY: cleavage-site(Smal)	
(B) LOCATION: 5257	
(C) IDENTIFICATION METHOD: S	
(ix) FEATURE:	
(A) NAME/KEY: cleavage-site(Kpnl)	
(B) LOCATION: 23952400	
(C) IDENTIFICATION METHOD: S	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
CTCGCTTTCC TTGTTCCTGA CCGATAACAT CACTGAGATC ATGTTGTAGC GCCCGGGATA	60
CTGCATCAGT TGGTTTCGGG CGTTCGAGAG CGTGCTTACC TTCCAGAAAC GCACAGACAG	120
CTTGCAGATG ATCGCCTATC AGGCATCCTT CACCGTTAAT TAGCCCCACT TCATCTTCGT	180
TATCTTTCGC GACGATAATT TTTCTGCCCG ACTTAATAGC TTCAGTTGCA CTGGAGATTG	240
CGCCGGGAAC GCCACGCAGA GCGCCTGTAA GCGCCAGTTC TCCGACTAAT TCATATTCAT	300
CTAACTTATT GGCTGTAAGC TGTTCTGAGG CCGCCAGCAA CGCAATGGCG ATAGGTAAAT	360
CATATCGTCC CCCTTCTTTT GGCAGATCAG CTGGAGCCAG GTTGATGGTG ATTTTTTTCG	420
CCGGATATTC ATATCCGCTA TTGATAATGG CGCTGCGCAC GCGATCGCGA GCTTCTTTTA	480

	CONTRICTO TOUTABLE ACCATES ITA ASCESSORIAS ACCITIACIO ATATOTACO.	040
	CAACAGTGAT CGGGGGCGCA TTTACTCCCA GGGCTGCGCG GGTATGAACA ATTGACAGTG	600
	ACATAAGCCC TCCTTGAGTC ACCATTATGT GCATAAGATA TCGCTGCTGT AGCCCGCTAA	660
	TTCGTGAATT TTAGTGGCTG ATTCCTGTTT ATTTGTGCAA GTGAAGTTGA GTTGTTCTGC	720
	CGGTGGAATG ATGCTCGCAA AAATGCAGCG GACAAAGGAT GAACTACGAG GAAGGGAACA	780
	ACATTCATAC TGAAATTGAA TTTTTTTCAC TCACTATTTT ATTTTTAAAA AACAACAATT	840
	TATATTGAAA TTATTAAACG CATCATAAAA ATCGGCCAAA AAATATCTTG TACTATTTAC	900
	AAAACCTATG GTAACTCTTT AGGCATTCCT TCGAACAAGA TGCAAGAAAA GACAAA	956
	ATG ACA GCC CTT CTA CGA GTG ATT AGC CTG GTC GTG ATT AGC GTG GTG	1004
	Met Thr Ala Leu Leu Arg Val Ile Ser Leu Val Val Ile Ser Val Val	
	1 5 10 15	
	GTG ATT ATT ATC CCA CCG TGC GGG GCT GCA CTT GGA CGA GGA AAG GCT	1052
	Val Ile Ile Ile Pro Pro Cys Gly Ala Ala Leu Gly Arg Gly Lys Ala	
	20 25 30	
	TAGAGATCAA GCCTTAACGA ACTAAGACCC CCGCACCGAA AGGTCCGGGG GTTTTTTTTTC	1112
i	ACCTTAAAAA CATAACCGAG GAGCAGACAA TGAATAACAG CACAAAATTC TGTTTCTCAA	1172
	GATTCAGGAC GGGGAACTÁA CT ATG AAT GGC GCA CAG TGG GTG GTA CAT GCG	1224
	Met Asn Gly Ala Gln Trp Val Val His Ala	
	1 5 10	
)	TTG CGG GCA CAG GGT GTG AAC ACC GTT TTC GGT TAT CCG GGT GGC GCA	1272
	Leu Arg Ala Gln Gly Val Asn Thr Val Phe Gly Tyr Pro Gly Gly Ala	
	15 20 25	
5	ATT ATG CCG GTT TAC GAT GCA TTG TAT GAC GGC GGC GTG GAG CAC TTG	1320
	Ile Met Pro Val Tyr Asp Ala Leu Tyr Asp Gly Gly Val Glu His Leu	
	30 35 40	
)	CTA TGC CGA CAT GAG CAG GGT GCG GCA ATG GCG GCT ATC GGT TAT GCT	1368
	Leu Cys Arg His Glu Gln Gly Ala Ala Met Ala Ala Ile Gly Tyr Ala	
	45 50 55	
	CGT GCT ACC GGC AAA ACT GGC GTA TGT ATC GCC ACG TCT GGT CCG GGC	1416
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	GCA ACC AAC CTG ATA ACC GGG CTT. GCG GAC GCA CTG TTA GAT TCC ATC	1464
)	Ala Thr Asn Leu Ile Thr Gly Leu Ala Asp Ala Leu Leu Asp Ser Ile	
	75 80 85 90	

	CCT	GTT	GTT	GCC	ATC	ACC	GGT	CAA	GTG	TCC	GCA	CCG	TTT	ATC	GGC	ACT	1512
	Pro	Val	Val	Ala	Ile	Thr	Gly	Gln	Val	Ser	Ala	Pro	Phe	Ile	Gly	Thr	
5		•			95					100					105		
	GAC	GCA	TTT	CAG	GAA	GTG	GAT	GTC	CTG	GGA	TTG	TCG	TTA	GCC	TGT	ACC	1560
	Asp	Ala	Phe	Gln	Glu	Val	Asp	Val	Leu	Gly	Leu	Ser	Leu	Ala	Cys	Thr	
10				110					115					120			
	AAG	CAT	AGC	TTT	CTG	GTG	CAG	TCG	CTG	GAA	GAG	TTG	CCG	CGC	ATC	ATG	1608
	Lys	His	Ser	Phe	Leu	Val	Gln	Ser	Leu	G1u	Glu	Leu	Pro	Arg	Ile	Met	
			125					130					135			•	
15	GCT	GAA	GCA	TTC	GAC	GTT	GCC	TGC	TCA	GGT	CGT	CCT	GGT	CCG	GTT	CTG	1656
	Ala	Glu	Ala	Phe	Asp	Val	Ala	Cys	Ser	Gly	Arg	Pro	Gly	Pro	Val	Leu	
		140					145					150					
20	GTC	GAT	ATC	CCA	AAA	GAT	ATC	CAG	TTA	GCC	AGC	GGT	GAC	CTG	GAA	CCG	1704
20	Val	Asp	Ile	Pro	Lys	Asp	Ile	Gln	Leu	Ala	Ser	Gly	Asp	Leu	Glu	Pro	
	155					160					165					170	
	TGG	TTC	ACC	ACC	GTT	GAA	AAC	GAA	GTG	ACT	TTC	CCA	CAT	GCC	GAA	GTT	1752
25	Trp	Phe	Thr	Thr	Val	Glu	Asn	Glu	Val	Thr	Phe	Pro	His	Ala	Glu	Val	
					175					180					185		
	GAG	CAA	GCG	CGC	CAG	ATG	CTG	GCA	AAA	GCG	CAA	AAA	CCG	ATG	CTG	TAC	1800
30	Glu	Gln	Ala	Arg	Gln	Met	Leu	Ala	Lys	Ala	Gln	Lys	Pro	Met	Leu	Tyr	
00				190					195					200			
	GTT	GGC	GGT	GGC	GTG	GGT	ATG	GCG	CAG	GCA	GTT	CCG	GCT	TTG	CGT	GAA	1848
	Val	Gly	Gly	Gly	Val	Gly	Met	Ala	Gln	Ala	Val	Pro	Ala	Leu	Arg	Glu	
35			205					210					215				
	TTT	CTC	GCT	GCC	ACA	AAA	ATG	CCT	GCC	ACC	TGT	ACG	CTG	AAA	GGG	CTG	1896
	Phe	Leu	Ala	Ala	Thr	Lys	Met	Pro	Ala	Thr	Cys	Thr	Leu	Lys	Gly	Leu	
40		220					225					230					
	GGC	GCA	GTA	GAA	GCA	GAT	TAT	CCG	TAC	TAT	CTG	GGC	ATG	CTG	GGG	ATG	1944
	Gly	Ala	Val	Glu	Ala	Asp	Tyr	Pro	Tyr	Tyr	Leu	Gly	Met	Leu	Gly	Met	
	235					240					245					250	
45	CAC	GGC	ACC	AAA	GCG	GCA	AAC	TTC	GCG	GTG	CAG	GAG	TGT	GAC	CTG	CTG	1992
	His	Gly	Thr	Lys	Ala	Ala	Asn	Phe	Ala	Val	Gln	Glu	Cys	Asp	Leu	Leu	
					255					260					265		
50	ATC	GCC	GTG	GGC	GCA	CGT	TTT	GAT	GAC	CGG	GTG	ACC	GGC	AAA	CTG	AAC	2040
	Ile	Ala	Val	Gly	Ala	Arg	Phe	Asp	Asp	Arg	Val	Thr	Gly	Lys	Leu	Asn	
				270					275					280			

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Glu Met Asn Lys Leu Arg Gln Ala His Val Ala Leu Gln Gly Asp Leu  300 305 310  AAT GCT CTG TTA CCA GCA TTA CAG CAG CCG TTA AAT CAA TGT GAC TGG Asn Ala Leu Leu Pro Ala Leu Gln Gln Pro Leu Asn Gln Cys Asp Trp 315 320 325 330  CAG CAA CAC TGC GCG CAG CTG CGT GAT GAA CAT TCC TGG CGT TAC GAC Gln Gln His Cys Ala Gln Leu Arg Asp Glu His Ser Trp Arg Tyr Asp 335 340 345  CAT CCC GGT GAC GCT ATC TAC GCG CGG TTG TTG TTG TTA AAA CAA CTG TCG His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Lys Gln Leu Ser 350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTC ACC ACA GAT GTG GGG CAG CAC Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365 370 375  CAG ATG TGG GCT GCG CAG CAC CAC ACC GCC CCG CAC CCC GCG GAA AAT TTC 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ACC GGT TTT GCT TTA CCG GCG GCG 350 ASP ATG ACC TCC AGC GGT TA GGT ACC ACA GAT GTT GCT TTA CCG GCG GCG ATC GCC GAA GCC GCG CAC CAC ACC ACC GCT TTA CCG GCG GCG 41e Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 405  GTT GGC GCA CAA GTC GCG CAC CCG ACC GAT ACC GTT GTC TTA CCC GCG GCG 410 Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 410 GTT GGC GCA CAC GCT TTC ATG ATG ATG GTG CAC GCT GGC CAC CTA AAA 410 GTT GGC GCA CAC GTT TTC ATG ATG ATG GTG CAC CAC GAT ACC GTG GGC ACC GTA AAA 420 425  GGT GAC GCC TTT TC ATG ATG ATG GTG CTTA CTC GAT AAC CAA CCG TTA AAA 420 425  GGT GAC GCA TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 420 425  GGT GAC GCA TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 425  GGT GAC GCA TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 426  GGT GAC GCA TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 427  428  430 435  440  450  450  450  450  450  450		ACC	TTC	GCG	CCA	CAC	GCC	AGT	GTT	ATC	CAT	ATG	GAT	ATC	GAC	CCG	GCA	2088
GAA ATG AAC AAG CTG CGT CAG GCA CAT GTG GCA TTA CAA GGT GAT TTA  Glu Met Asn Lys Leu Arg Gln Ala His Val Ala Leu Gln Gly Asp Leu  300 305 310  AAT GCT CTG TTA CCA GCA TTA CAG CAG CCG TTA AAT CAA TGT GAC TGG Asn Ala Leu Leu Pro Ala Leu Gln Gln Pro Leu Asn Gln Cys Asp Trp  315 320 325 330  75 CAG CAA CAC TGC GCG CAG CTG CGT GAT GAA CAT TCC TGG CGT TAC GAC Gln Gln His Cys Ala Gln Leu Arg Asp Glu His Ser Trp Arg Tyr Asp  335 340 345  CAT CCC GGT GAC GCT ATC TAC GCG CGG TTG TTG TTA AAA CAA CTG TCC His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser  350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTC ACC ACA GAT GTG GGG CAC CAC  Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His  365 370 375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC  380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG CGC  11e Tr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  395 400 405  ATC ACC TCC AGC GAT GCG CGA CCG ACC ACC GGT TTT GGT TTA CCG GCG CGC  11e Tr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  396 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ACC GAT GGT TTT GGT TTA CCG GCG GCG  11e Tr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  397 400 405  GTT GGG GCA CAA GTC GCG CAC ACC GCC AAC GAT GCC GCC GTA AAA  398 400 405  GTT GGC GCA CAA GTC GCG CAC ACC GAC GAT ACC GTT GTC TGT ATC TCC  410 GTT GGC GCA CAA GTC GCG CGA ACC GAC GAT ACC GTT GTC TGT ATC TCC  410 GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  410 GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  411 GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  412 GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  415 GGC GAT GAT CCC TTT CT ATG ATG ATC GTC TTA CTC GAT AAC CAA CGG TTA AAC  410 GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  410 GTT GGC GCA CAA GTC GCG CTC TTT CTT ATC GTC TTA CTC GAT AAC CAA CGG TTA AAC  410 GTT GGC GCA CAA GTC GCG CTC TTT TTT ATT CTC GAT AAC CAA CGG TTA AAC  420 425  430 430 430 430 440  44		Thr	Phe	Ala	Pro	His	Ala	Ser	Val	Ile	His	Met	Asp	11	Asp	Pro	Ala	
Glu Met Asn Lys Leu Arg Gln Ala His Val Ala Leu Gln Gly Asp Leu 300 305 310  AAT GCT CTG TTA CCA GCA TTA CAG CAG CCG TTA AAT CAA TGT GAC TGG Asn Ala Leu Leu Pro Ala Leu Gln Gln Pro Leu Asn Gln Cys Asp Trp 315 320 325 330  TAG CAA CAC TGC GCG CAG CTG CGT GAT GAA CAT TCC TGG CGT TAC GAC Gln Gln His Cys Ala Gln Leu Arg Asp Glu His Ser Trp Arg Tyr Asp 335 340 345  CAT CCC GGT GAC GCT ATC TAC GCG CG CTG TTG TTG TTA AAA CAA CTG TCG His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser 350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTG ACC ACA GAT GTG GGG CAG CAC Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365 370 375  CAG ATG TGC GCG GCT GCG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ACA GTT TTT GGT TTA CCG GCG GCG 11e Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GCA CAA GTC GCG CAG CCA ACT GCC CTG TTT GTT GTT ATC TCC 415 Asp Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GAC TCT TC ATG ATG AAT GTG CAA GAG CTG GCG CAC CTA AAA 430 435  GGT GAC GAC TTT CCA TTT AAA ATC GTC TTA CTC GAT AAA CAC CTG TA CAA 430 435  GGG ATC GTT CGA CAA TTT CGT TTA AAA CAA CTG CTG CAA CAA CTG GTG GTG ACC ACC ATC GCC CTG AAA 430 435  GGG ATC GTT CCA CAC TTT CCA TTT AAT CTT CTT AAT CTC CAAA CAA C	5			285					290					295				
AAT GCT CTG TTA CCA GCA TTA CAG CAG CCG TTA AAT CAA TGT GAC TGG 21 Asn Ala Leu Leu Pro Ala Leu Gln Gln Pro Leu Asn Gln Cys Asp Trp 315 320 325 330  15 CAG CAA CAC TGC GCG CAG CTG CGT GAT GAA CAT TCC TGG CGT TAC GAC Gln Gln His Cys Ala Gln Leu Arg Asp Glu His Ser Trp Arg Tyr Asp 335 340 345  CAT CCC GGT GAC GCT ATC TAC GCG CGC CTG TTG TTG TTA AAA CAA CTG TCG His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser 350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTC GTC ACC ACA GAT GTG GGG CAG CAC Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365 370 375  CAG ATG TGC GCG CAG CAC ATC GCC CAC ATC GCC CAC CAC CCG GAA AAT TTC 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ACC ACT CGC CCG GAA AAT TTC 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ACC ACC GTT TTT GGT TTA CCG GCG GCG 11e Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GAC ACA GTC GCG CAG CAC ACC GCG AAC GTC GTC TGT ACC GCT GTC TGT ACC GCT GTC GTC GTC GTC GTC GTC GTC GTC G		GAA	ATG	AAC	AAG	CTG	CGT	CAG	GCA	CAT	GTG	GCA	TTA	CAA	GGT	GAT	TTA	2136
AAT GCT CTG TTA CCA GCA TTA CAG CAG CCG TTA AAT CAA TGT GAC TGG  Asn Ala Leu Leu Pro Ala Leu Gln Gln Pro Leu Asn Gln Cys Asp Trp  315  320  325  330  330  325  330  330  33		Glu	Met	Asn	Lys	Leu	Arg	Gln	Ala	His	Val	Ala	Leu	Gln	Gly	Asp	Leu	
Asn Ala Leu Leu Pro Ala Leu Gln Gln Pro Leu Asn Gln Cys Asp Trp 315  CAG CAA CAC TGC GCG CAG CTG CGT GAT GAA CAT TCC TGG CGT TAC GAC Gln Gln His Cys Ala Gln Leu Arg Asp Glu His Ser Trp Arg Tyr Asp 335  CAT CCC GGT GAC GCT ATC TAC GCG CCG TTG TTG TTA AAA CAA CTG TCG His Pro Gly Asp Ala Ille Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser 350  GAT CGT AAA CCT GCG GAT TGC GTG ACC ACA GAT GTG GGG CAG CAC Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365  GAT GG GCT GCG CAG CAC ATC GCC CAC ACT GCC CAC ACT CGC CCG GAA AAT TTC 360  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG ATC GTG GGC GCA CAC GCG CAC CAC ACC GCT GTC GTC TTA CTC ATG GGT GAC GCC CAC ACT GGC GAT ACC ATG GGT TTT GGT TTA CCG GCG GCG ATC GTG GGC GCA CAC ATC GCC CAC ACT GGT GTC TTA CTC GCC CAC ATC GCC GCA CAC ATC GCC CAC ACT GGT TTA CTC GCC CCG ATC ACC TCC AGC GCG CCG CCG CCG ACC GAT ACC ATC GCC CCG GCG GCG ATC ACC TCC AGC GCG CCG CCG CCG ACC GAT ACC ATC GCC CCG GCG GCG ATC GTT GGC GCA CAC ATC GCC CAC ACC GAT ACC GTT GTC TTA CTC CCC ACC ACC ACC GCA CCG ACC GAA GAC GAT ACC GTT GTC TTC CCC CCG ACC GCA CCG ACC GCA CCG ACC GAT ACC GTT GTC TCC CCC ACC GCG GCG CCG CCG ACC GAA GAG GTT GCC GCC ACC GTA AAA ATC GTC GCC GCC CCC CCC CCC CCC ACC GTC GCC ACC GTA AAA ATC GTC GCC GCC CCC ACC GAA GAG CTG GCC ACC GTA AAA ATC GTC GCC GCC CCC CCC CCC CCC ACC GTA AAA ATC GTC GCC GCC CCC CCC CCC ACC GAT ACC ACC GTA AAA ATC GTC GCC GCC CCC CCC CCC ACC GTA AAA ACC GCC TTA AAA ATC GTC GCC CCC CCC CCC CCC CCC ACC GTA AAA ATC GTC GTC GTC GTC GTC GTC GTC GTC GTC G	10		300					305					310					
315   320   325   330   330   345   330   346   345   346   345   346   345   346   345   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346   346		AAT	GCT	CTG	TTA	CCA	GCA	TTA	CAG	CAG	CCG	TTA	AAT	CAA	TGT	GAC	TGG	2184
CAG CAA CAC TGC GCG CAG CTG CGT GAT GAA CAT TCC TGG CGT TAC GAC  Gln Gln His Cys Ala Gln Leu Arg Asp Glu His Ser Trp Arg Tyr Asp  335  CAT CCC GGT GAC GCT ATC TAC GCG CCG TTG TTG TTG TTA AAA CAA CTG TCG  His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser  350  GAT CGT AAA CCT GCG GAT TGC GTC GTG ACC ACA GAT GTG GGG CAG CAC  Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His  365  CAG ATG TGG GCT GAC CAC ATC GCC CAC ACT CGC CAG GAA AAT TTC  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe  380  ATC ACC TCC AGG GGT TTA GGT ACC ATG GGT TTT GGT TTA CG GCG GCG  ATC ACC TCC AGG GGT TTA GGT ACC ATG GGT TTT GGT TTA CG GCG GCG  ATG TGG GCA CAA GTC GCG CAG CAC ACT GGT TTA CG GCG GCG  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CG GCG GCG  ATC GGC GCA CAA GTC GCG CAA CAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415  420  425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CGT ACC GCA CAC CAC ACC GCG ACC ACC GTA AAA  256  GGT GAC GCC TCT TTC ATG ATG AAT GTG CAA GAG CTG GCC ACC GTA AAA  257  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GCC ACC GTA AAA  257  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GCC ACC GTA AAA  258  GGG AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  410  421  422  425  426  GGT GAC GAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  427  428  429  420  425  426  GGT GAC GAC GAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  427  438  440  450  455  GGG AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  450  450  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC CAA CGC  GTY Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		Asn	Ala	Leu	Leu	Pro	Ala	Leu	Gln	Gln	Pro	Leu	Asn	G1n	Cys	Asp	Trp	
GIN GIN HIS CYS Ala GIN Leu Arg Asp Glu HIS Ser Trp Arg Tyr Asp 335 340 345  CAT CCC GGT GAC GCT ATC TAC TAC GCG CCG TTG TTG TTA AAA CAA CTG TCG His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Leu Leu Ser 350 355 360  GAT CGT AAA CCT GCG GAT TGC TGC GTG ACC ACA GAT GTG GGG CAG CAC Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly GIn His 365 370 375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CAG AAA TTTC 23  GIN Met Trp Ala Ala GIn His Ile Ala His Thr Arg Pro Glu Asn Phe 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG 11e Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GAC AAA GTC GCG CGA CAG ACC GAT ACC GTT GTC TGT ATC TCC Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGT ACC GAC GTA AAA 25  GIY Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  45 CGC AAG CAG CTA CAG CAG TTG CAG CAG CAG CTG TTT TTT CAG GAA CGA TAG CAG CAG CAG CGG CGG CGG CGG CGG CGG C		315					320					325					330	
20 CAT CCC GGT GAC GCT ATC TAC GCG CCG TTG TTG TTA AAA CAA CTG TCG 21 His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Leu Lys Gln Leu Ser 350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTG ACC ACA GAT GTG GGG CAG CAC 23 Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365 370 375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC 23 Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG 11e Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GCA CAA GTC GCG CAG CAC AAC GAT ACC GTT GTC TGT ATC TCC 41 GGT GAC GGT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 410  GTT GGC GCA CAA GTC GCG CAG CCG AAC GAT ACC GTT GTC TGT ATC TCC 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 415  GGT GAC GGC TTA CATG ATG AAT GTG CAA GAG CTG GGT AAC CAA CAC GAT CAC 426  GGT GAC GGC TTA CATG ATG AAT GTG CAA GAG CTG GAT AAC CAA CAC CAA CAC 427  428  429  420  425  GGT GAC GAC CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CAC CAA CAC GAT ACC 440  450  450  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CAC CAA CAC CAC CAC ATG ATG CAC CAC GTT ATC CAC CAC CAC CAC CAC CAC CAC CAC C	15	CAG	CAA	CAC	TGC	GCG	CAG	CTG	CGT	GAT	GAA	CAT	TCC	TGG	CGT	TAC	GAC	2232
CAT CCC GGT GAC GCT ATC TAC GCG CCG TTG TTG TTA AAA CAA CTG TCG His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser 350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTG ACC ACA GAT GTG GGG CAG CAC Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC 370  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe 380  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395  GGT GGC GCA CAA GTC GCG CGA ACC GAT ACC GTT GTC TGT ATC TCC Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415  GGT GAC GGC TCT TC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  GGT GAC GGC TCT TC AGG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 430  430  435  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC CAA CGG TTA 440  45  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 45  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC GIY Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		Gln	Gln	His	Cys	Ala	Gln	Leu	Arg	Asp	Glu	His	Ser	Trp	Arg	Tyr	Asp	
His Pro Gly Asp Ala Ile Tyr Ala Pro Leu Leu Leu Lys Gln Leu Ser  350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTG ACC ACA GAT GTG GGG CAG CAC 23 Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His  365 370 375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC 23 Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe  380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG 11e Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 401 GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC 42 Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25 Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430 435 436  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 25 Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser						335					340					345		
350 355 360  GAT CGT AAA CCT GCG GAT TGC GTC GTC ACC ACA GAT GTG GGG CAG CAC 23  Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365 370 375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC 23  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG 24  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC 24  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GGC TCT TTC ATG ATG ATG GTC GAA GAG CTG GGC ACC GTA AAA 25  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  45 CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA AGG 25  GGG ATG GTT CGA CAA TGG CAG CAG CTG TTT TTT CAG GAA CGA TAC AGC 26  GGG ATG GTT CGA CAA TGG CAG CAG CTG TTT TTT CAG GAA CGA TAC AGC 26  GGG ATG GTT CGA CAA TGG CAG CAG CTG TTT TTT CAG GAA CGA TAC AGC 26  GIy Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	20	CAT	CCC	GGT	GAC	GCT	ATC	TAC	GCG	CCG	TTG	TTG	TTA	AAA	CAA	CTG	TCG	2280
GAT CGT AAA CCT GCG GAT TGC GTC GTC ACC ACA GAT GTG GGG CAG CAC  Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His  365  370  375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC  23  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe  380  385  390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  395  400  405  405  410  GTT GGC GCA CAA GTC GCG CGA ACC GAT ACC GTT GTC TGT ATC TCC  24  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415  420  425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA  25  GGY ASP Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430  435  436  GGC AAG CAG CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC CAG CTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		His	Pro	Gly	Asp	Ala	Ile	Tyr	Ala	Pro	Leu	Leu	Leu	Lys	Gln	Leu	Ser	
Asp Arg Lys Pro Ala Asp Cys Val Val Thr Thr Asp Val Gly Gln His 365 370 375  CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC 23  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG 1le Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC 415  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  GGY Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser					350					355					360			
CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe  380  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  395  400  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA  25  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA  45  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		GAT	CGT	AAA	CCT	GCG	GAT	TGC	GTC	GTG	ACC	ACA	GAT	GTG	GGG	CAG	CAC	2328
CAG ATG TGG GCT GCG CAG CAC ATC GCC CAC ACT CGC CCG GAA AAT TTC  Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe  380  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  35  395  400  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415  420  425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430  435  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	?5	Asp	Arg		Pro	Ala	Asp	Cys	Val	Val	Thr	Thr	Asp	Val	Gly	Gln	His	
Gln Met Trp Ala Ala Gln His Ile Ala His Thr Arg Pro Glu Asn Phe 380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  45 CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser				365					370					375				
380 385 390  ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  395 400 405 410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430 435 440  45 CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 25  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC 26  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		CAG	ATG	TGG	GCT	GCG	CAG	CAC	ATC	GCC	CAC	ACT	CGC	CCG	GAA	AAT	TTC	2376
ATC ACC TCC AGC GGT TTA GGT ACC ATG GGT TTT GGT TTA CCG GCG GCG  Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala  395  400  405  410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415  420  425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA  25  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430  435  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	30	Gln	Met	Trp	Ala	Ala	Gln	His	Ile	Ala	His	Thr	Arg	Pro	Glu	Asn	Phe	
Ile Thr Ser Ser Gly Leu Gly Thr Met Gly Phe Gly Leu Pro Ala Ala 395 400 405 410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  GCG AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 25  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC 26  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser			380					385					390					
395 400 405 410  GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430 435 440  45 CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		ATC	ACC	TCC	AGC	GGT	TTA	GGT	ACC	ATG	GGT	TTT	GGT	TTA	CCG	GCG	GCG	2424
GTT GGC GCA CAA GTC GCG CGA CCG AAC GAT ACC GTT GTC TGT ATC TCC  Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser  415  420  425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA  Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys  430  435  440  45  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		Ile	Thr	Ser	Ser	G1y	Leu	Gly	Thr	Met	Gly	Phe	Gly	Leu	Pro	Ala	Ala	
Val Gly Ala Gln Val Ala Arg Pro Asn Asp Thr Val Val Cys Ile Ser 415 420 425  GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25 Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	<b>35</b> .																	
GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25 Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		GTT	GGC	GCA	CAA	GTC	GCG	CGA	CCG	AAC	GAT	ACC	GTT	GTC	TGT	ATC	TCC	2472
GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25 Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  GGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		Val	Gly	Ala	Gln	Val	Ala	Arg	Pro	Asn	Asp	Thr	Val	Val	Cys		Ser	
GGT GAC GGC TCT TTC ATG ATG AAT GTG CAA GAG CTG GGC ACC GTA AAA 25 Gly Asp Gly Ser Phe Met Met Asn Val Gln Glu Leu Gly Thr Val Lys 430 435 440  45 CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	40																	
430 435 440  CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA 25  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser																		2520
CGC AAG CAG TTA CCG TTG AAA ATC GTC TTA CTC GAT AAC CAA CGG TTA  Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu  445  450  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		Gly	Asp	Gly	Ser	Phe	Met	Met	Asn		Gln	Glu	Leu	Gly	Thr	Val	Lys	
Arg Lys Gln Leu Pro Leu Lys Ile Val Leu Leu Asp Asn Gln Arg Leu 445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser						•												
445 450 455  GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC 26  Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	<b>4</b> 5	CGC	AAG	CAG	TTA	CCG	TTG	AAA	ATC	GTC	TTA	CTC	GAT	AAC	CAA	CGG	TTA	2568
GGG ATG GTT CGA CAA TGG CAG CAA CTG TTT TTT CAG GAA CGA TAC AGC Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser		Arg	Lys	Gln	Leu	Pro	Leu	Lys	Ile	Val	Leu	Leu	Asp	Asn	Gln	Arg	Leu	
Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser				445					450					455				
Gly Met Val Arg Gln Trp Gln Gln Leu Phe Phe Gln Glu Arg Tyr Ser	50	GGG	ATG	GTT	CGA	CAA	TGG	CAG	CAA	CTG	TTT	TTT	CAG	GAA	CGA	TAC	AGC	2616
460 465 470	-	Gly	Met	Val	Arg	Gln	Trp	Gln	Gln	Leu	Phe	Phe	Gln	Glu	Arg	Tyr	Ser	
			460					465					470					

	GAA	ACC	ACC	CTT	ACT	GAT	AAC	CCC	GAT	TTC	CTC	ATG	TTA	GCC	AGC	GCC	2664
	Glu	Thr	Thr	Leu	Thr	Asp	Asn	Pro	Asp	Phe	Leu	Met	Leu	Ala	Ser	Ala	
5	475	•			•	480					485					490	
	TTC	GGC	ATC	CAT	GGC	CAA	CAC	ATC	ACC	CGG	AAA	GAC	CAG	GTT	GAA	GCG	2712
	Phe	Gly	Ile	His	Gly	Gln	His	Ile	Thr	Arg	Lys	Asp	Gln	Val	Glu	Ala	
10					495					500					505		
	GCA	CTC	GAC	ACC	ATG	CTG	AAC	AGT	GAT	GGG	CCA	TAC	CTG	CTT	CAT	GTC	2760
	Ala	Leu	Asp	Thr	Met	Leu	Asn	Ser	Asp	Gly	Pro	Tyr	Leu	Leu	His	Val	
				510					515					520			
15	TCA	ATC	GAC	GAA	CTT	GAG	AAC	GTC	TGG	CCG	CTG	GTG	CCG	CCT	GGC	GCC	2808
	Ser	Ile	Asp	Glu	Leu	Glu	Asn	Val	Trp	Pro	Leu	Val	Pro	Pro	Gly	Ala	
			525					530					535				
20	AGT	AAT	TCA	GAA	ATG	TTG	GAG	AAA	TTA	TCA	TGA						2841
	Ser	Asn	Ser	Glu	Met	Leu	Glu	Lys	Leu	Ser							
		540					545					•					
25																	
23	(2)	INFO	DRMA1	LION	FOR	SEQ	ID !	10:2	:								
		(i)	SEC	QUENC	CE C	iara(	CTER	ISTIC	cs:								
			(/	A) LE	ENGTI	1: 54	18 an	nino	acio	is							
30	-		(F	3) TY	PE:	amir	o ac	çid									
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				LECUL													
35						-				D NO							
	_	Asn	Gly	Ala		Trp	Val	Val	His		Leu	Arg	Ala	Gln		Val	
	I				5	_	_			10			_		15		
	Asn	Thr	Val		Gly	Tyr	Pro	Gly		Ala	He	Met	Pro	Val	Tyr	Asp	
40				20					25					30			
	Ala	Leu		Asp	Gly	Gly	Val	Glu	His	Leu	Leu	Cys	Arg	His	Glu	Gln	
			35		_			40					45				
45	Gly	Ala	Ala	Met	Ala	Ala		Gly	Tyr	Ala	Arg		Thr	Gly	Lys	Thr	
		50	_				55					60					
		Val	Cys	lle	Ala		Ser	G1 y	Pro	Gly		Thr	Asn	Leu	He		
50	65					70					75					80	
	Gly	Leu	Ala	Asp		Leu	Leu	Asp	Ser		Pro	Val	Val	Ala		Thr	
					85					90					95		

Gly	Gln	Val	Ser	Ala	Pro	Phe	Ile	Gly	Thr	Asp	Ala	Phe	Gln	Glu	Val
			100					105					110		
Asp	Val	Leu	Gly	Leu	Ser	Leu	Ala	Cys	Thr	Lys	His	Ser	Phe	Leu	Val
		115					120					125			
G1n	Ser	Leu	Glu	Glu	Leu	Pro	Arg	Ile	Met	Ala	Glu	Ala	Phe	Asp	Val
	130					135					140				
Ala	Cys	Ser	Gly	Arg	Pro	Gly	Pro	Val	Leu	Val	Asp	Ile	Pro	Lys	Asp
145					150					155					160
Ile	Gln	Leu	Ala	Ser	Gly	Asp	Leu	Glu	Pro	Trp	Phe	Thr	Thr	Val	Glu
				165					170					175	
Asn	Glu	Val	Thr	Phe	Pro	His	Ala	Glu	Val	Glu	G1n	Ala	Arg	Gln	Met
			180					185					190		
Leu	Ala	Lys	Ala	Gln	Lys	Pro	Met	Leu	Tyr	Val	Gly	Gly	Gly	Val	Gly
		195					200					205			
Met	Ala	Gln	Ala	Val	Pro	Ala	Leu	Arg	Glu	Phe	Leu	Ala	Ala	Thr	Lys
	210					215					220				
Met	Pro	Ala	Thr	Cys	Thr	Leu	Lys	Gly	Leu	Gly	Ala	Val	Glu	Ala	Asp.
225					230					235					240
Tyr	Pro	Tyr	Tyr	Leu	Gly	Met	Leu	Gly	Met	His	Gly	Thr	Lys	Ala	Ala
				245					250					255	
Asn	Phe	Ala	Val	Gln	Glu	Cys	Asp	Leu	Leu	Ile	Ala	Val	Gly	Ala	Arg
			260					265					270		
Phe	Asp	Asp	Arg	Val	Thr	Gly	Lys	Leu	Asn	Thr	Phe	Ala	Pro	His	Ala
		275	~				280					285			
Ser	Val	Ile	His	Met	Asp	Ile	Asp	Pro	Ala	Glu	Met	Asn	Lys	Leu	Arg
	290					295					300				
Gln	Ala	His	Val	Ala	Leu	Gln	Gly	Asp	Leu	Asn	Ala	Leu	Leu	Pro	Ala
305					310					315					320
Leu	Gln	Gln	Pro	Leu	Asn	Gln	Cys	Asp	Trp	Gln	Gln	His	Cys	Ala	Gln
				325					330					335	
Leu	Arg	Asp	Glu	His	Ser	Trp	Arg	Tyr	Asp	His	Pro	Gly	Asp	Ala	He
			340					345					350		
Tyr	Ala	Pro	Leu	Leu	Leu	Lys	Gln	Leu	Ser	Asp	Arg	L.ys	Pro	Ala	Asp
		355					360					365			
Cys	Val	Val	Thr	Thr	Asp	Val	Gly	Gln	His	Gln	Met	Trp	Ala	Ala	Gln
	370					375					380				

His Ile Ala His Thr Arg Pro Glu Asn Phe Ile Thr Ser Ser Gly Leu

	385					390					395					400	
5	Gly	Thr	Met	Gly	Phe	Gly	Leu	Pro	Ala	Ala	Val	Gly	Ala	Gln	Val	Ala	
					405					410					415		
	Arg	Pro	Asn	Asp	Thr	Val	Val	Cys	Ile	Ser	Gly	Asp	Gly	Ser	Phe	Met	
10				420					425					430			
	Met	Asn	Val	Gln	Glu	Leu	Gly	Thr	Val	Lys	Arg	Lys	Gln	Leu	Pro	Leu	
			435					440					445				
15	Lys	Ile	Val	Leu	Leu	Asp	Asn	Gln	Arg	Leu	Gly	Met	Val	Arg	Gln	Trp	
		450					455					460					
	Gln	Gln	Leu	Phe	Phe	Gln	Glu	Arg	Tyr	Ser	Glu	Thr	Thr	Leu	Thr	Asp	
	465					470					475					480	-
20	Asn	Pro	Asp	Phe	Leu	Met	Leu	Ala	Ser	Ala	Phe	Gly	lle	His	G1y	Gln	
					485					490					495		
	His	He	Thr		Lys	Asp	Gln	Val		Ala	Ala	Leu	Asp	Thr	Met	Leu	
25				500					505					510			
	Asn	Ser		Gly	Pro	Tyr	Leu		His	Val	Ser	lle		Glu	Leu	Glu	
			515	_	_		_	520			_		525				
30	Asn		Trp	Pro	Leu	Val		Pro	Gly	Ala	Ser		Ser	Glu	Met	Leu	
	0.1	530		_			535					540.					
		Lys	Leu	Ser													
	545																
35	(2)	TATE	DMAT	r t O N T	EOD	CEA	TD. A	in · o ·									
	(4)				FOR CE Ch					•							
		(1)			ENGTI												
40					ZNOTI PE:												
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	,				POLO				,								
45		(i i)			LAR 1				nuc le	eic a	acid			-			
		(,			ESCRI							DNA*	•				
		(i i i)			ETICA				/								
50					ENSE:												
J.					CE DE		PTIO	ON: S	SEQ I	D NO	):3:			·			
	TAAC				CATO												22
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55																	

١.	(2) INFOR	MATION FOR SEQ ID NO:4:	
	(i) S	SEQUENCE CHARACTERISTICS:	
5		(A) LENGTH: 21 base pairs	
		(B) TYPE: nucleic acid	
		(C) STRANDEDNESS: single	
10		(D) TOPOLOGY: linear	
70	(ii) l	MOLECULAR TYPE: other nucleic acid	
		(A) DESCRIPTION: /desc="synthetic DNA"	
	(iii) l	HYPOTHETICAL: NO	
15	(iv) <i>i</i>	ANTI-SENSE: YES	
	(xi) 5	SEQUENCE DESCRIPTION: SEQ ID NO:4:	,
	TCTTTTCTTC	G CATCTTGTTC G	21
20	(2) INFORM	MATION FOR SEQ ID NO:5:	
	(i) S	SEQUENCE CHARACTERISTICS:	
		(A) LENGTH: 22 base pairs	
25		(B) TYPE: nucleic acid	
		(C) STRANDEDNESS: single	
		(D) TOPOLOGY: linear	
	(ii) N	MOLECULAR TYPE: other nucleic acid	
30		(A) DESCRIPTION: /desc="synthetic DNA"	
	(iii) H	HYPOTHETICAL: NO	
	(iv) A	ANTI-SENSE: NO	
35	(xi) S	SEQUENCE DESCRIPTION: SEQ ID NO:5:	
	TCTGTTTCTC	C AAGATTCAGG AC	22
	(2) INFORM	MATION FOR SEQ ID NO:6:	
40	(i) S	SEQUENCE CHARACTERISTICS:	
		(A) LENGTH: 19 base pairs	. •
		(B) TYPE: nucleic acid	
45		(C) STRANDEDNESS: single	
		(D) TOPOLOGY: linear	
	(ii) N	MOLECULAR TYPE: other nucleic acid	
		(A) DESCRIPTION: /desc="synthetic DNA"	
50	(iii) l	HYPOTHETICAL: NO	
	(iv) A	ANTI-SENSE: YES	
		•	

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6: CGCCGGTAAA CCAAAACCC

19

## 10 Claims

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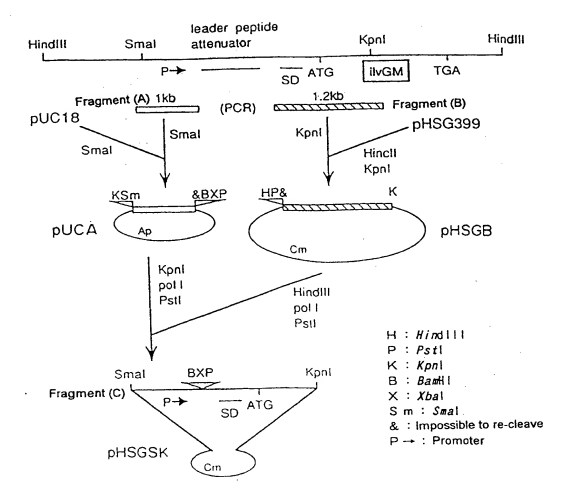
30

- 1. A microorganism belonging to the genus <u>Escherichia</u> and having a capability of producing L-valine or L-leucine, which requires lipoic acid for growth.
- 75 2. A microorganism belonging to the genus <u>Escherichia</u> and having a capability of producing L-valine or L-leucine, which is deficient in H<sup>+</sup>-ATPase activity.
  - 3. A microorganism belonging to the genus <u>Escherichia</u> and having a capability of producing L-valine or L-leucine, which requires lipoic acid for growth and which is deficient in H\*-ATPase activity.
  - 4. A microorganism according to any one of claims 1 to 3, which has the capability of producing L-valine by carrying a gene for L-valine biosynthesis, of which regulatory mechanism is substantially released.
- 5. A microorganism according to any one of claims 1 to 3, which has the capability of producing L-leucine by carrying a gene for L-leucine biosynthesis, of which regulatory mechanism is substantially released.
  - 6. A microorganism according to claim 4, which has the capability of producing L-valine by introducing a DNA fragment including an <u>ilvGMEDA</u> operon which express each of <u>ilvG</u>, <u>ilvM</u>, <u>ilvE</u> and <u>ilvD</u> genes, and does not express threonine deaminase, into a cell.
  - A microorganism according to claim 6 wherein a region of the <u>ilvGMEDA</u> operon necessary for attenuation by L-valine and/or L-isoleucine and/or L-leucine is deleted.
- 8. A microorganism according to claim 7 wherein the region necessary for the attenuation which is deleted has nucleotides 953 to 1160 of a sequence shown in SEQ ID NO: 1.
  - 9. A microorganism according to any one of claims 1 to 8, which is Escherichia coli.
  - 10. A microorganism according to claim 9, which is <u>Escherichia coli</u> W1485atpA401/pMWdAR6, W1485lip2/pMWdAR6 or AJ12631/pMWdAR6.
  - 11. A method for producing L-valine which comprises culturing the microorganism having the capability of producing L-valine as defined in any one of claims 1 to 4 and 6 to 9 in a liquid medium to allow L-valine to be produced and accumulated in the medium, and collecting it.
  - 12. A method for producing L-leucine which comprises culturing the microorganism having the capability of producing L-leucine as defined in any one of claims 1 to 3, 5 and 9 in a liquid medium to allow L-leucine to be produced and accumulated in the medium, and collecting it.

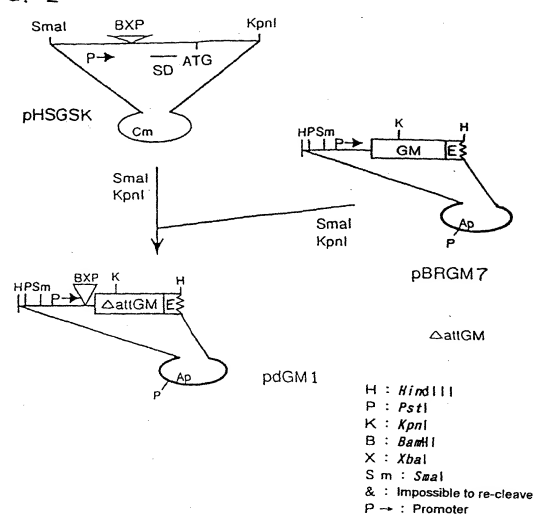
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FIG. 1



F | G. 2



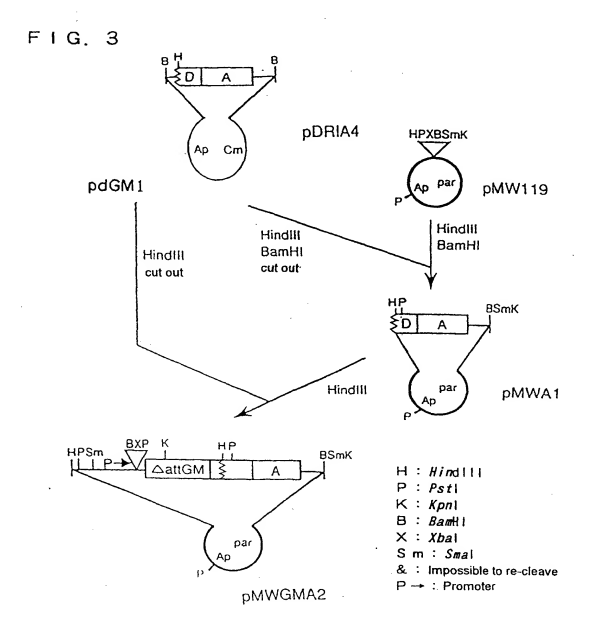
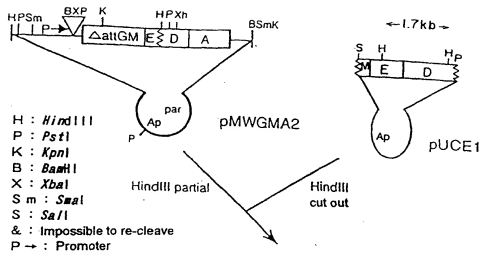


FIG. 4



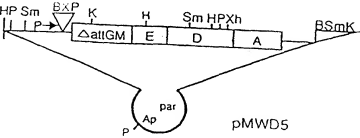


FIG. 5

# pMWD5 13kb



H: Hindlll

P : Pstl

B : BanHI

A : Accili

B g : Bg/1

SnaB I S : SnaBl Acc III partial N : Not1 X : Xbal

& : Impossible to re-cleave

ilvGMEDA\* Ap par

Self-ligation

pMWdAR6 11.7kb

## INTERNATIONAL SEARCH REPORT

International application No.

			PCT/J	P95/U1/19
Int.	SSIFICATION OF SUBJECT MATTER C1 <sup>6</sup> C12N1/21, C12N1/20, C C12R1:19), (C12P13/08	, Cl2P1:19)		(C12N1/21,
According to	o International Patent Classification (IPC) or to both		and IPC	
B. FIEL	DS SEARCHED			
	C16 C12N1/00-3/00, C12P13			
	on searched other than minimum documentation to the ex			
_	ta base committed during the international search (name of IS PREVIEWS, WPI	of data base and, where p	racticable, search t	erms used)
C. DOCU	MENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where ap	ppropriate, of the releva	nt passages	Relevant to claim No.
X A	JP, 5-137568, A (Ajinomoto June 1, 1993 (01. 06. 93)	CO., Inc.) (Family: none	, e)	1 - 3 4 - 12
х	HERBERT A A et al. "Lipoid Escherichia-coli and Other Arch. Microbiol., (1975(RI No. 3, p. 259-266	r Microorgan:	isms",	1, 3
А Р, Х	YOKOTA A, et al. "Pyrubic an F-1-ATPase-defective m			2, 4-12 1, 3
Р, А	coli W1485lip2", Biosciend Biochemistry, (1994. Dec) p. 2164-2167			2 4-12
F, A				2, 4-3.2
Р, Х	YOKOTA A, et al. "Pyrubic lipoic acid auxotroph of 1 W1485", Applied Microbiol Vol. 41, No. 6, p. 638-641	Escherichia ( . Biotech.,	coli	1, 3
X Furthe	er documents are listed in the continuation of Box C.	See patent f	family annex.	
"A" docume	Categories of cited documents: an defining the general state of the art which is not considered particular relevance	date and not in co		rnational filing date or priority cation but cited to understand invention
"E" earlier o	becamest but published on or after the international filing date ast which may throw doubts on priority chains(s) or which is establish the publication date of another citation or other	considered novel step when the do	or cannot be considered along the constant in	
"O" docume measu	reason (as specified) ent referring to an oral disclosure, une, exhibition or other	"Y" document of part considered to in combined with an being obvious to	volve sa inventive	cisimed invention cannot be step when the document is documents, such combination he art
	ent published prior to the international filing date but later than ority date claimed	"&" document membe	•	
i.	actual completion of the international search imber 21, 1995 (21. 11. 95)	Date of mailing of the December		(19. 12. 95)
Name and r	nailing address of the ISA/	Authorized officer	<del></del> -	
Japa	nese Patent Office			
Facsimile N	io.	Telephone No.		

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# INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP95/01719

-1	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
ategory*	Change of accomment, with indication, where appropriate,		
, а			2, 4-12
A	Hiroshi Aida and four others "Amino Acid Fermentation" May 30, 1986 (30. 05. 86) Gakkai Shuppan Center, p. 397-422		1 - 12
<b>. A</b> .	<pre>JP, 4-330275, A (Ajinomoto Co., Inc.), November 18, 1992 (18. 11. 92) (Family: none)</pre>		1 - 12
A	<pre>JP, 61-185195, A (Mitsui Toatsu Chemicals, Inc.), August 18, 1986 (18. 08. 86) (Family: none)</pre>		1 1.2
A	EP, 519113, Al (Ajinomoto K.K.), December 23, 1992 (23. 12. 92) (Family: none)		1 1.2
A	WO, 8702984, A (Amer. Biogenetics Co.), May 21, 1987 (21. 05. 87) & AU, 8767372, A & EP, 245497, A & JP, 63501687, W		1 - 12
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